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- © Color photographs, a process for preparing them and color photographic materials employed therefor.
- There is provided a color photograph improved in preservability that has been obtained by making chemically inactive the aromatic amine type color developing agents and their oxidized product that remains in the silver halide photographic materials after color development processing. The color photograph exhibits excellent performance in that its white background can be prevented from discoloring even during long-term storage or display, and deterioration of a dye image caused by the remaining color developing agent being taken into the photographic material after the color development, bleaching, and fixing processes, or due to its Noxidized product, can be prevented.

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COLOR PHOTOGRAPHS, A PROCESS FOR PREPARING THEM AND COLOR PHOTOGRAPHIC MATERIALS EMPLOYED THEREFOR

BACKGROUND OF THE INVENTION

(1) Field of the Invention

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The present invention relates to color photographs improved in preservability, and particularly to color photographs improved in preservability that have been obtained by making chemically inactive the aromatic amine type color developing agents (hereinafter referred to as aromatic amine developing agents) and their oxidized product that remain in the silver halide photographic materials after color development processing, and to a process of the production of said color photographs and silver halide color photographic materials employed therefor.

(2) Description of the Prior Art

In the field of silver halide color photographic materials, for example, as dye image forming couplers (hereinafter referred to as couplers), there have been developed, along with couplers that give bright cyan, magenta, and yellow dyes with less subsidiary absorption that afford good color reproduction, highly active couplers through which color development completes within a short time. New additives and other agents to draw further enhance the excellent performance of these couplers are also being developed. However, in actually this new performance caused a deterioration of the preservability of the color photographs due to interaction with the components of the processing solutions that remain in the photosensitive material after the processing.

It is known that, of the processing solution components remaining in the photographic material after the development processing, in particular the aromatic primary amine compound, that is, a developing agent and the compounds derived therefrom, damages the fastness of the image under the influence, for example, of light, heat, and oxygen during long-term storage, or they themselves cause self-coupling or interact with coexistents to change to colored materials, resulting in so-called "stain". This can be considered a fatal defect in a color photograph.

On the other hand, many studies have been made to prevent images from deteriorating and to prevent stain. For example, ideas have been suggested to employ couplers that fade out less, to use fading preventive agents to prevent fading due to light, or to use ultraviolet absorbing agents to prevent an image from being deteriorated by ultraviolet rays.

Although it is recognized that the above compounds have an effect as agents to prevent a dye image from fading or discoloring, the compounds cannot successfully meet the customer demand for high quality images, and they have not yet achieved overall excellence due to their problems of changing the hue, causing fogging or defective dispersion, or forming fine crystals after the application of the emulsion.

However, the stain in question in the invention is produced when aromatic amine compounds taken into the color photograph after the development processing are oxidized with oxygen or the like during prolonged storage of the color photograph, and which at the same time react with contained colorless compounds, such as color image forming compounds (couplers), to form colored compounds. The inventors of this invention tried to develop methods for scavenging aromatic amine compounds taken into the color photograph or the oxidized product of such aromatic amine compounds. However, since the effect of these scavenging compounds also often decreased, for example due to decomposition or deterioration during long-term storage of the color photograph, and the amount of the aromatic amine compounds taken into the color photograph changed notably with the type of development processing, in actual fact the compounds intended to scavenge either the involved aromatic amine compounds or the oxidized product thereof had been unsatisfactory.

BRIEF SUMMARY OF THE INVENTION

Therefore an object of the invention is to provide a process of the production of a color photograph wherein the white background is prevented from discoloring during long-term storage or display after colordevelopment, bleaching, and fixing of the silver halide color photographic material.

Another object of the present invention is to provide a color photograph which is prevented from deterioration of the dye image due to a remaining color developing agent taken into the photographic material after the color development, bleaching, and fixing.

Still another object of the invention is to provide a process of forming a color image of a color photographic material wherein even if the color photographic material is processed with a processing solution in a running state, a processing solution that will be washed with less water or will not be washed with water, a processing solution that is substantially free from benzyl alcohol, such as a color developing solution, and whose components will be brought into the photographic material in a greater amount, or other processing solutions that will, for example, impose a burden on the color development, image deterioration due to the remaining aromatic amine developing agent or its oxidized product and the occurrence of stain or the side effects therefrom can be prevented.

Other and further objects, features and advantages of the invention will appear more fully from the following description.

DETAILED DESCRIPTION OF THE INVENTION

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The inventors have studied intensively to attain the above objects by tracing accurately aromatic amines that have been brought into the color photograph during the photographic processing and the resultant reaction species which change with time and by quickly making inactive the aromatic amines or the subsequent reaction species using scavenging compounds relevant to the reaction species, which have resulting in the present invention.

Thus the present invention provides (1) a color photograph improved in preservability that contains both a compound (A) that can combine chemically with the aromatic amine developing agent remaining after the color development processing to produce a chemically inactive and substantially colorless compound, and a compound (B) that can combine chemically with the oxidized product of the aromatic amine developing agent remaining after the color development processing to produce a chemically inactive and substantially colorless compound, (2) a process of the production of a color photograph improved in preservability by processing a silver halide color photographic material in the presence of compound (A) and compound (B), and (3) a silver halide color photographic material that contains compound (A) and compound (B) in at least one layer of the hydrophilic colloid layers on a base of the silver halide color photographic material.

In the specification and claims, "chemically inactive compound" means (1) a compound that does not or hardly decompose chemically for a long period of time or (2) a compound that does not promote the fading of the dye, does not generate colored materials by the acceleration of the decomposition of the residual coupler, or does not form colored materials, even if it decomposes. In the specification and claims, "substantially colorless compound" means (1) a compound that has no absorption at the visible ray range longer than 350 nm (2) a compound that has a molecular extinction coefficient of 1,000 or below at the visible ray range longer than 350 nm or (3) a compound that gives a color photograph having a white background with reflection density (optical density) of 0.01 or below in respect of a yellow, magenta or cyan dye.

Methods of allowing the preservative compound (A) and the preservative compound (B) to coexist in a color photograph (e.g., a color print and a color film) obtained by processing a silver halide color photographic material to attain the objects of the present invention include:

1) a method wherein at least one of compounds (A) and (B) is previously contained during a step of the production of the photographic material, and compound (A) and/or compound (B) are contained in one or more layers of the hydrophilic colloid layers on the base, and if both compound (A) and compound (B) are contained they may be contained in the same layer or different layers, with the former being preferable. The hydrophilic colloid layers include photosensitive layers and non-photosensitive layers, such as silver halide emulsion layers, ultraviolet absorbing layers, and protective layers.

2) a method wherein before, during, or after the color development processing the photographic material is processed with a processing solution to which compound (A) and/or compound (B) has been added, to allow compound (A) and/or compound (B) to be contained in the color photograph,

which compound (A) may essentially be added to a color developing solution, but compound (A) is preferably added to a processing solution after the development processing with a view to avoiding the reaction of compound (A) with the color developing agent in the color developing solution, and

3) a method wherein one of compounds (A) and (B) is contained in a step of the production of the photographic material, and the other compound is added to any processing solution used before, during, or after the color development processing.

The aromatic amine developing agents in this specification and claim herein include aromatic primary, secondary, and tertiary amine compounds, and more particularly phenylenediamine type compounds and aminophenol type compounds. Typical examples thereof are 3-methyl-4-amino-N,N-diethylaniline, 3-methyl-4-amino-N-ethyl-N- β -hydroxyethylaniline, 3-methyl-4-amino-N-ethyl-N- β -methanesulfonamidoethylaniline, 4-methyl-2-amino-N,N-diethylaniline, 4-methyl-2-amino-N-ethyl-N- β -methanesulfonamidoethylaniline, 2-amino-N-ethyl-N- β -hydroxylethylaniline, 3-methyl-4-methylamino-N-ethyl-N- β -methanesulfonamidoethylaniline, 3-methyl-4-butylamino-N,N-diethylaniline, 3-methyl-4-acetylamino-N-ethyl-N- β -methanesulfonamidoethylaniline, 3-methyl-4-methylaniline, 3-methyl-4-methylaniline, 3-methyl-4-methylaniline, 3-methyl-4-benzylamino-N-ethyl-N- β -methanesulfonamidoethylaniline, and 3-methyl-4-cyclohexylamino-N-ethyl-N-methylaniline, and their sulfates, hydrochlorides, phosphates, or p-toluenesulfonates, tetraphenyl borates, p-(t-octyl)benzenesulfonates, o-aminophenol, p-aminophenol, 4-amino-2-methylphenol, 2-amino-3-methylphenol, and 2-oxy-3-amino-1,4-dimethylbenzene.

As other examples can be mentioned those described in L.F.A. Mason, "Photographic Processing Chemistry" Focal Press (1966), pages 226 to 229, U.S.Patent Nos. 2,193,015 and 2,592,364, and Japanese Patent Application (OPI) No. 64933/1973.

The oxidized products of the aromatic amine developing compounds in this specification and claims means oxidized products derived chemically by removing one or two electrons from the aromatic amine developing agents.

Of the compounds (A) that can chemically bond with the aromatic amine developing agent after the color development processing to form substantially colorless compounds, preferable ones are compounds that can react with the rate constant k_2 (at 80°C) of the secondary reaction with p-anisidine within the range of 1.0 t/mol.sec to 1 x 10 5 t /mol.sec.

If the k_2 is too great the compounds themselves become unstable and react with gelatin and water to decompose. On the other hand, if the k_2 is too small the reaction of the compounds with the aromatic amine developing agents is slow, and as a result the side effect of the remaining aromatic amine developing agents that the invention intends to obviate cannot be prevented.

Of such compounds (A), preferable ones can be represented by the following general formula (I) or (II): General formula (I)

General formula (II)

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wherein R₁ and R₂ each represent an aliphatic group (preferably one having 1~60 carbon atoms, more preferably having 10~50 carbon atoms), an aromatic group (preferably one having 6~60 carbon atoms, more preferably having 16~50 carbon atoms), or a heterocyclic group (preferably one having 2~60 carbon atoms). X represents a group that can react with the aromatic amine developing agent to split off, A represents a group that can react with the aromatic amine developing agent to form a chemical bond, n is 1 or 0, B represents a hydrogen atom, an aliphatic group (preferably one having 1~60 carbon atoms,more preferably having 10~50 carbon atoms), an aromatic group (preferably one having 6~60 carbon atoms,more preferably having 16~50 carbon atoms), a heterocyclic group (preferably one having 2~60 carbon atoms,more preferably having 10~50 carbon atoms), or an acyl or a sulfonyl group (preferably one having 1~50 carbon atoms,more preferably one having 2~50 carbon atoms), Y represents a group that can facilitate the addition of the aromatic amine developing agent to the compound having general formula (II), and R₁ and X together or Y and R₂ or B together may combine to form a ring structure.

Of ways wherein the remaining aromatic amine developing agent and the compound (A) chemically combine, typical ways are substitution reactions and addition reactions.

Groups of the compounds represented by general formulae (I) and (II) are described further.

The aliphatic groups represented by R_1 , R_2 and B may be straight chain, branched chain or cyclic alkyl groups, alkenyl group or alkynyl groups that may be substituted. The aromatic groups represented by R_1 , R_2 and B may be any of the carbocyclic aromatic group (e.g., phenyl and naphtyl), and the heterocyclic

aromatic group (e.g., furyl, thienyl, pyrazolyl, pyridyl and indolyl), may be of a monocyclic type or a condensed ring type (e.g., benzofuryl and phenanthridinyl). These groups may be further substituted.

The heterocyclic groups represented by R1 , R2 and B are preferably groups having a 3-to 10membered ring structure comprising carbon atoms, oxygen atoms, nitrogen atoms, sulfur atoms, or hydrogen atoms, and the hetero ring may itself be a saturated or unsaturated ring, and may be substituted (e.g., chromanyl, pyrrolidinyl, pyrrolinyl, and morpholinyl).

X represents a group that can react with the aromatic amine developing agent to split off, and preferably represents a group that attaches to A via an oxygen atom, a sulfur atom, a nitrogen atom (e.g., 2pyridyloxy, 2-pyrimidyloxy, 4-pyrimidyloxy, 2-(1,2,3-triazine)oxy, 2-benzimidazolyl, 2-imidazolyl, 2-thiazolyl, 2-benzthlazolyl, 2-furyloxy, 2-thiophenyloxy, 4-pyridyloxy, 3-isooxazolyloxy, 3-pyrazolidinyloxy, 3-oxo-2pyrazolonyl, 2-oxo-1-pyridinyl, 4-oxo-1-pyridinyl, 1-benzimidazolyl, 3-pyrazolyloxy, 3H-1,2,4-oxadiazolin-5oxy, aryloxy, alkoxy, alkylthio, arylthio, and substituted N-oxy), or a halogen atom.

A represents a group that can react with the aromatic amine developing agent to form a chemical bond, and it includes a group containing a low electron density atom such as

When X is a halogen atom, n is 0. L represents a single bond, an alkylene group, -O-, -S-,

(e.g., carbonyl, sulfonyl, sulfinyl, oxycarbonyl, phosphonyl, thiocarbonyl, aminocarbonyl and silyloxy).

Y has the same meaning as that of Y in general formula (II), and Y' has the same meaning as that of Y. R' and R" may be the same or different, and each represents -L"-Ro.

 R_0 has the same meaning as that of R_1 . $R^{\star \star}$ represents a hydrogen atom, an aliphatic group (e.g., methyl, isobutyl, t-butyl, vinyl, benzyl, octadecyl, and cyclohexyl), an aromatic group (e.g., a phenyl, pyridyl, and naphthyl), a heterocyclic group (e.g., piperidinyl, pyranyl, furanyl, and chromanyl), an acyl group (e.g., acetyl, and benzoyl), or a sulfonyl group (e.g., methanesulfonyl and benzenesulfonyl).

In particular, A represents preferably a divalent O O O II group represented by -O- C -, -S- C -or -alkylene- C -.

Preferred compounds of those represented by the general formula (I) are those represented by general formula (I-a), (I-b), (I-c) or (I-d) that can react with the rate constant K2 (at 80°C) of the secondary reaction with p-anisidine within the range of 1 x 10 1 to 1 x 10 5 L /mol.sec.

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$$(I-a)$$

$$R_1$$
-Link-C-O-Ar

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(I-b)

(I-c) $R_1-Link-C-O-C$ N

In the above formulae, R₁ has the same meaning as R₁ in general formula (I); Link represents a single bond or -O-; Ar represents an aromatic group having the same meanings as defined in R₁, R₂ and B, provided that the group released as a result of reaction with an aromatic amine developing agent is not a group useful as a photographic reducing agent such as catechol derivative. Ra, Rb and Rc, which may be the same or different, each represent a hydrogen atom, an aliphatic, aromatic or heterocyclic group having the same meaning as defined in R1, R2 and B. Further, Ra, Rb and Rc each represent an alkoxy group, aryloxy group, heterocyclooxy group, alkylthio group, arylthio group, heterocyclothio group, amino group, alkylamino group, acyl group, amido group, sulfonamide group, sulfonyl group, alkoxycarbonyl group, sulfo group, carboxyl group, hydroxyl group, acyloxy group, ureido group, urthane group, carbamoyl group or sulfamoyl group. Ra and Rb, or Rb and Rc may combine toghther to form a 5-to 7-membered heterocyclic ring which may be further substituted by a substituent, may form, a spirocyclic ring or bicyclo ring, or may be condensed by an aromatic ring. Z₁ and Z₂ each represent a non-metal atom group necessary to form a 5-to 7-membered heterocyclic ring which may be further substituted by a substituent, may form a spirocyclic ring or bicyclo ring, or may be condensed by an aromatic ring. The compound released as a result of the reaction of Z₁ with an aromatic amine developing agent is not a coupler or 1-phenyl-3pyrazolidones.

The adjustment of the rate constant k_2 (at 80°C) of the secondary reaction with p-anisidine within the range 1 x 10 1 to 1 x 10 51 /mol.sec for the compounds represented by general formulae (l-a)~(l-d), especially for the compound represented general formula (l-a), can be attained by selecting a substituent, when Ar is a cyclocarbon aromatic group. In this case, the sum total of Hammett's σ -value of substituents, which may be dependent on the kind of group of R₁, is preferably 0.2 or greater, more preferably 0.4 or greater, most preferably 0.6 or greater.

It is preferable that the sum total of carbon atoms of the compound is more than 13, when a compound represented by the general formula (I-a) to (I-b) is added to produce a photographic material. It is not desirable that these compounds decompose during the development processing, in order to achieve the object of the invention.

Y in general formula (II) is preferably an oxygen atom, a sulfur atom, = N-R4 or



Herein, R₄, R₆ and R₆ each represent a hydrogen atom, an aliphatic group (preferably having 1~30 carbon atoms,more preferably having 1~20 carbon atoms, e.g., methyl, Isopropyl, t-butyl, vinyl, benzyl, octadecyl and cyclohexyl), an aromatic group (preferably having 6~40 carbon atoms,more preferably having 6~30 carbon atoms, e.g., phenyl, pyridyl, and naphthyl), a heterocyclic group (preferably having 2~30 carbon atoms,more preferably having 2~20 carbon atoms, e.g., piperidyl, pyranyl, furanyl and chromanyl), an acyl group (preferably having 2~30 carbon atoms,more preferably having 2~20 carbon atoms, e.g., acetyl and benzoyl), or a sulfonyl group (preferably having 1~30 carbon atoms,more preferably having 1~20 carbon atoms, e.g., methanesulfonyl, and benzenesulfonyl), and R₆ and R₆ may bond together to form a ring structure.

Of the compounds represented by general formulae (I) and (II), the compounds of general formula (I) are especially preferable. In these compounds, the compound represented by general formula (I-a) or (I-c) is more preferable, and the former is more preferable.

Typical examples of these compounds are given below, but the invention is not limited to these compounds.

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(I-2)

$$\begin{array}{c|c}
N-O \\
\downarrow & O \\
N-O & \downarrow & (n)
\end{array}$$

(I-3)

$$\begin{array}{c}
N-0 \\
N = 0 \\
0 \\
OCOC_7 H_{15}(n)
\end{array}$$

(1-4)

$$(t)_{C_5H_{11}} - \underbrace{\begin{array}{c} O & O \\ O & \parallel \\ OCH_2CH_2CH_2CH_2C-O-N \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O & O \\ \parallel \\ O & H \end{array}}_{C_5H_{11}(t)} + \underbrace{\begin{array}{c} O$$

$$\begin{array}{c} (t)_{C_5H_{11}} - \begin{array}{c} O \\ -O - CH_2CH_2CH_2CH_2C - O - N \\ C_5H_{11}(t) \end{array}$$

$$\begin{array}{c}
\text{(t)} & \text{O} \\
\text{C}_{5}\text{H}_{11} - \text{O} - \text{CH}_{2}\text{CH}_{2}\text{CH}_{2}\text{C} - \text{O} - \text{N} \\
\text{C}_{5}\text{H}_{11}(t) & \text{O}
\end{array}$$

$$\begin{array}{c|c}
C_2H_5 \\
C_5H_{11} & OCHC-O-N
\end{array}$$

$$\begin{array}{c}
(I - 8) \\
C_{2}H_{5} \\
0 \\
C_{5}H_{11}
\end{array}$$

$$\begin{array}{c}
C_{2}H_{5} \\
0 \\
C_{5}H_{11}(t)
\end{array}$$

$$\begin{array}{c}
C_{4}H_{9}CHOCS \\
C_{2}H_{5}
\end{array}$$

(I - / 0)

$$\begin{array}{c|c}
C_4 H_9 CHOCS \\
C_2 H_5
\end{array}$$

$$\begin{array}{c|c}
C_5 H_{11}(t) \\
C_5 H_{11}(t)
\end{array}$$

(1 - / /)

CONHCH₂CH₂CH₂CH₂O
$$C_5H_{11}^{(t)}$$

40 (1-/2)

(n) C₁₃H₂₇CC

OCH₃

5

15

$$(I - / 3)$$

$$\begin{array}{cccc} & & & & C_2H_5 & O \\ \text{(n)} & & & & \parallel \\ & & & C_4H_9\,\text{CHCH}_2\,\text{OCO}. \end{array}$$

$$(1 - / s)$$

$$(1 - 19)$$

$$(1-20)$$

$$C_{12}H_{25}$$
 $SO_2CH_2CH_2B_1$

10

15 (I-22)

(1-23)

O
$$C_7H_{15}OCO - SO_2 - OH$$

(I - 2 ¢)

$$C_7H_{15}OCO-N$$

(I - 25)

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5 (I-26)

$$C_{2}H_{5}$$
 O $C_{4}H_{9}CHCH_{2}O-C-O$ N N $OC_{16}H_{33}$

25 CH₃ O CH₃ C₂H₅ CH₂OC CHC₄H₅

$$C_{13}H_{27}C-O$$
 N
 N
 $C_{2}H_{5}$
 $C_{5}H_{11}(t)$

(I-30)

$$C_2H_5$$
 $C_4H_9CHCH_2O$
 $OCH_2CHC_4H_9$
 $OCH_2CHC_4H_9$

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(1-32)

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(1-33)

(I-34)

$$C_{4}H_{9}-CH-O-C-O CH_{3} CH_{2}O-C NHCOC_{13}H_{27}$$

(I-35)

C₂H₅ O CH₃

$$C_4H_9 - CHCH_2O - C - O$$

$$N$$

$$N$$

$$N$$

$$OC H$$

(1-36)

5

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25

(I-37)

(1-38)

$$C_{10}H_{21}-O-C-O-$$

(I-39)

⁴⁵ (I-40)

(I-41)

([-42)

$$C_{2}H_{5}$$
 O $C(CH_{3})_{3}$
 $C_{4}H_{9}-CHCH_{2}O-C-O-(N-C(CH_{3})_{3})$

(1-43)

$$C_{2}H_{5}$$
 O O $C_{4}H_{9}$ - CHCH₂-O-C-O-C-C(CH₃)₃

-50

$$\begin{array}{c|c}
H & CH_3 \\
\hline
 & C_2H_5 \\
\hline
 & OC-OCH_2CHC_4H_9 (n) \\
\hline
 & OC-OCH_2CHC_4H_9 (n)
\end{array}$$

$$\begin{array}{c|c}
OCH_2CHC_4H_9 (n) \\
\hline
 & OCH_2CHC_4H_9 (n)
\end{array}$$

(I-45)

(I-46)

(I-47)

$$\begin{array}{c} C_2H_5 & O \\ (n)C_4H_9 - CHCH_2O - CO \\ N & OC - CH_2CH - C_4H_9 (n) \\ Cl & Cl O \end{array}$$

$$(1-48)$$

$$(I-49)$$

(I-51)

$$\begin{array}{c|c}
C = CHSO_2 - \left(\begin{array}{c} \\ \\ \end{array}\right) \\
O - C - O - CH_2 CH - C_4 H_9 (n) \\
0 & C_2 H_5
\end{array}$$

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(1-52)

(n)
$$C_7H_{15}$$
 $-O$ $-C_7H_{15}$ (n) C_7H_{15} $-O$ $-C_7H_{15}$ (n) O

²⁰ (I-53)

$$\begin{array}{c} C_2H_5 \\ \text{(a)} C_4H_9 CHCH_2-O-CO- \\ \text{O} \\ \end{array} \begin{array}{c} C_2 \\ \text{OC} \\ \text{OC} \\ \text{OC} \\ \text{OC} \\ \text{OC} \\ \text{C}_2H_5 \\ \text{OC} \\ \text{OC} \\ \text{C}_1H_2 \\ \text{C}_2H_2 \\ \text{C}_1H_2 \\ \text{C}_2H_2 \\ \text{C}_$$

(I−54)

(I-55)

(1-56)

$$(t)C_{5}H_{11}- \underbrace{\begin{array}{c} C_{2}H_{5} & O & C\ell \\ O & C_{2}H_{5} \\ O - CHCH_{2}O - CO - CO - CH_{2}CHO - CH_{2}CHO - CO - CH_{2}CHO - CO - CH_{2}CHO - CO - CH_{2}CHO - CH_{2}CHO - CO - CH_{2}CHO - CH_{2}CHO - CO - CH_{2}CHO - CO - CH_{2}CHO - CO - CH_{2}CHO - CH_{2}CHO - CO - CH_{2}CHO -$$

(1-57)

$$(n)C_{15}H_{31}-CO \xrightarrow{Br} SO_2 \xrightarrow{Br} OC-C_{15}H_{31}(n)$$

(1-58)

$$\text{(n)} C_{15} H_{31} - \underset{0}{\overset{CQ}{\longleftarrow}} SO_2 - \underset{CQ}{\overset{CQ}{\longleftarrow}} OC - C_{15} H_{31} \text{(n)}$$

(1-59)

$$(1-60)$$

(I-61)

•

(I-64)

$$\begin{array}{c|c}
O & C_2H_5 \\
O-C-OCH_2CH-C_4H_9(n)
\end{array}$$

$$\begin{array}{c|c}
Cl & Cl \\
CO_2C_2H_5
\end{array}$$

(I-69)

CO₂C₂H₅

$$\begin{array}{c}
O \\
O - C - C_{15} H_{31} (n) \\
C \ell \\
\downarrow
\end{array}$$

$$(I-74)$$

$$(I-75)$$

SO₂CH₃

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$$(1-77)$$

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$$Cl \xrightarrow{O \\ Cl} Cl CH_2CH_2S \xrightarrow{OC_4H_9} Cl_{C_8H_{17}(t)}$$

 $\begin{array}{c} O \\ \parallel \\ O-C-OCH_2CH_2NHCOC_{15}H_{31} \text{(n)} \end{array}$ CO₂C₂H₅

-CH₂CH₂SO₂-C₁₆H₃₃(n) CO₂C₂H₅

0

$$(I-82)$$

$$C\ell \xrightarrow{OC_{4}H_{9}^{(n)}} CC_{8}H_{17}^{(t)}$$

(1-84)

$$\begin{array}{c} S & O \\ O & O \\$$

$$(1-85)$$

$$\begin{array}{c|c} CH_3 & CH_3 \\ N & OSO_2 & CH_3 \\ \hline \\ C-OCH_2 CHC_4 H_9 \text{ (n)} \\ 0 & C_2 H_5 \end{array}$$

$$(1-86)$$

$$\begin{array}{c} O & C\ell \\ O & C$$

$$(1-87)$$

(1-88)

$$\begin{array}{c} O \\ O \\ \parallel \\ - \\ SO_2 \end{array} \begin{array}{c} C\ell \\ - \\ OSOC_4H_9 \end{array} (n)$$

25 ([-89)

(1 - 90)

$$\begin{array}{c|c}
(n) \\
C_{16}H_{33}OC \\
\hline
C_{2}
\end{array}$$

$$\begin{array}{c|c}
C_{16}H_{33}OC \\
\hline
C_{2}
\end{array}$$

$$\begin{array}{c|c}
C_{16}H_{33}OC \\
\hline
C_{2}
\end{array}$$

$$\begin{array}{c|c}
C_{16}H_{33}
\end{array}$$

$$\begin{array}{c|c}
C_{16}H_{33}
\end{array}$$

$$\begin{array}{c|c}
C_{16}H_{33}
\end{array}$$

0

Synthesis examples of representative compounds of the present invention will now be described.

Synthesis Example 1 (Synthesis of exemplified compound 1-9)

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Synthesis of 2-ethylhexyl 4-dodecylbenzenethiocarbonate (exemplified compound I-9)

150 mL of chloroform and 9.9 mL (0.071 mol) of tiethylamine were added to 18 g (0.065 mol) of 4-dodecylbenzenethiol to dissolve it, and the solution was stirred at 25°C. 13.3 g (0.068 mol) of 2-ethylhexyl chlorocarbonate was added to the solution dropwise. After stirring for 30 min, cold aqueous hydrochloric acid was added thereto. After separation, the resulting chloroform layer was washed three times with cold

water and then dried over Glauber's salt. The Glauber's salt was filtered out, and the chloroform was then removed by distillation. The product thus obtained was purified by column chromatography, thereby obtaining 17.2 g of exemplified compound I-9 as an oil in a yield of 61.2%. Results of elementary analysis ($C_{2l}H_{46}O_2S_2$):

C H S

Found (%) 74.34 10.66 14.91

Calculated (%) 74.60 10.67 14.75

Synthesis Example 2 (Synthesis of exemplified compound I-13)

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i) Synthesis of 5-(3-hexadecyloxyphenyl)-3-hydroxy-1-pyenylpyrazole

60 mL of toluene and 10 g (0.12 mol) of manganese dioxide were added to 6.3 g (0.013 mol) of 4,5-dihydroxy-5-(3-hexadecyloxyphenyl-3-hydroxy-1-phenylpyrazole, and the mixture was heated in a steam bath for 2 hours with stirring. Inorganic substances were filtered out. The filtrate thus obtained was evaporated to dryness, followed by crystallization from 20 mL of ethyl acetate, thereby obtaining 5.8 g of a product having a melting point of 108 to 109 °C in a yield of 92.5 %.

ii) Synthesis of 3-(2-ethylhexyloxycarbonyloxy)-5-(3-hexadecyloxypenyl)-1-phenyl-pyrazole (exemplified compound I-13)

50 mL of chloroform and 1.0 mL (0.014 mol) of triethylamine were added to 5.3 g (0.011 mol) of 5-(3-hexadecyloxyphenyl)-3-hydroxy-1-phenylpyrazole to dissolve it, and the solution was stirred at 25°C. 2.3 g (0.012 mol) of 2-ethylhexyl chlorocarbonate was added to the solution dropwise. After stirring for 30 min, cold water was added thereto, and the separated chloroform layer was washed twice with 50 mL of cold water and then dried over Glauber's salt. The Glauber's salt was filtered out, and the chloroform was then removed by distillation. The product thus obtained was purified by column chromatography to produce 5.7 g of exemplified compound I-13 as an oil in a yield of 82%. Results of elementary analysis (C₄₀H₅₀N₂O₄):

	Č	H	N
Found (%)	76.13	9.47	4.11
Calculated (%)	75.91	9.56	4.43

Synthesis Example 3 (Synthesis of exemplified compound I-24)

Synthesis of 4-heptyloxycarbonyloxypyridine (exemplified compound 1-24)

100 mL of chloroform and 7.3 mL (0.052 mol) of triethylamine were added to 4.5 g (0.040 mol) of 4-hydroxypyridine monohydrate to dissolve it, and the solution was stirred at 25°C. 8.9 g (0.050 mol) of heptyl chlorocarbonate was added to the solution dropwise. After stirring for 30 min, cold aqueous hydrochloric acid was added thereto. After separation the resulting chloroform layer was washed twice with cold water and then dried over Glauber's salt. After filtering out the Glauber's salt, the chloroform was

removed by distillation, and the obtained product was purified by column chromatography, followed by crystallization from ethanol, thereby obtaining 7.5 g of exemplified compound I-24 having a melting point of 44 to 50°C in a yield of 83 %.

Results of elementary analysis (CpHeNO₃):

		С	H	N
10	Found (%)	69.52	8.47	6.03
*	Calculated (%)	69.31	8.55	6.22

5 Synthesis Example 4 (Synthesis of exemplified compound I-54)

150 mt acetonitrile was added to 19.4 of 3,3',5,5'-tetrachloro-4,4'-dihydroxybiphenylsulfone and 16.8 g of triethylamine, followed by stirring. 21.1 g of 2-ethylhexyl chloroformate was added thereto dropwise at room temperature. The stirring was continued for 3 hours, followed by extraction with ethyl acetate. The ethyl acetate layer was washed with water and then dried. Then the dried ethyl acetate layer was concentrated and the concentrate was purified by silica gel column chromatography to produce 20.5 g (58.4%) of white crystalline exemplified compound I-54. The melting point of the compound was 65 to 66°C.

Results of elementary analysis (C30H36Ct4O6S):

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		С	H	C L	S
30	Found (%)	51.41	5.47	20.21	4.61
	Calculated (%)	51.44	5.47	20.24	4.58

5 Synthesis Example 5-1 (Synthesis of exemplified compound I-57)

300 mL of acetonitrile was added to 11.3 g of 3, 3,5,5 -tetrabromobiphenylsulfone and 6.1 mL of triethylamine, followed by stirring. 12.3 g palmitic acid chloride was added thereto dropwise at room temperature. After the stirring was continued for 5 hours, the reaction mixture was poured into 500 mL of water. The precipitated crystals were collected by filtration, washed with water and dried, followed by recrystallization from chloroform/ethyl acetate mixed solvent to produce 175 g (84.0%) of crystalline exemplified compound I-57. The melting point of the product was 125 to 126°C. Results of elementary analysis (C₄₄H₈₆Br₄O₆S)

	•	C.	Н	Br	S
	Found (%)	50.60	6.21	30.39	3.11
50	Calculated (%)	50.68	6.38	30.66	3.07

Synthesis Example 5-2 (Synthesis of exemplified compound I-70)

14.3 mt of triethylamine was added to a solution consisting of 23.1 g of ethyl 3,5-dichloro-4-hydroxybenzoate and 100 mt of acetonitrile, and the solution was stirred at room temperature. Then, 30 g of hexadecyl chloroformate was added to the solution dropwise. After stirring for 1 hours, the solution was

poured into ice-water, and the resulting crystals were separated by filtration. By recrystallization of crude crystal from isopropyl alcohol the desired comound I-70 was obtained as 43.5 g (yield 87.9%) of white crystal. melting point: 42-43°C

Results of elementary analysis (C25H40C12O5)

C H CL
Found (%) 62.11 7.98 14.02
Calculated (%) 62.01 8.01 14.08

The compounds (B) that can chemically combine with the oxidized product of the aromatic amine developing agent to form a substantially colorless compound are preferably those having a nucleophilic group derived from a nucleophilic functional group that have a Pearson s nucleophilic ⁿCH₃ I value [R.G. Pearson et al., J. Am. Chem. Soc., 90, 319(1968)] of 5 or more.

More preferable examples of the compound (B) are those represented by the following general formula (III): General formula (III)

R-Z-M

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wherein R₇ represents an aliphatic group (preferably one having 10~80 carbon atoms,more preferably having 20~60 carbon atoms), an aromatic group (preferably one having 16~86 carbon atoms,more preferably having 26~66 carbon atoms), or a heterocyclic group (preferably one having 12~82 carbon atoms,more preferably having 22~62 carbon atoms), Z represents a nucleophilic group, and M represents a hydrogen atom, a metal cation, an ammonium cation or a protective group.

The aliphatic group represented by R₇ is a straight chain, branched chain, or cyclic alkyl, alkenyl or alkynyl group. These group may be further substituted. The aromatic group represented by R₇ may be any of a carbocyclic aromatic group (e.g., phenyl, and naphthyl), and a heterocyclic aromatic group (e.g., furyl, thienyl, hydrazolyl, pyridyl, and indolyl), which may be of monocyclic type or condensed ring type (e.g., benzofuryl and phenanthridinyl). Further, these aromatic rings may have a substituent.

The heterocyclic group represented by R₂ is preferably one having a 3-to 10-membered ring structure comprising carbon atoms, oxygen atoms, nitrogen atoms, sulfur atoms, or hydrogen atoms. The heterocyclic ring itself may be a saturated ring or an unsaturated ring, and it may be substituted further with a substituent (e.g., chromanyl, pyrrolidyl, pyrrolinyl, and morpholinyl).

Z represents a nucleophilic group. The nucleophilic groups includes a group having an oxygen atom, a sulfur atom, or a nitrogen atom as an atom that will directly chemically combine with the oxidized product of the aromatic amine developing agent (Examples of the nucleophilic group include amine compounds, azide compounds, hydrazine compounds, mercapto compounds, sulfide compounds, sulfinic acid compounds, cyano compounds, thiocyano compounds, thiosulfuric acid compounds, seleno compounds, halide compounds, carboxy compounds, hydroxamic acid compounds, active methylene compounds, phenol compounds, and nitrogen heterocyclic compounds.

M represents a hydrogen atom, a metal cation, an ammonium cation or a protective group.

The compound represented by general formula (III) reacts with the oxidized product of the aromatic amine developing agent by a nucleophilic reaction (typically a coupling reaction).

Of compounds represented by general formula (III), the most preferable ones are those represented by the following general formula (IV):

General formula (IV)

 R_{13} R_{13} R_{12}

In the formula M' represents an atom or an atomic group that can form an inorganic salt (e.g., salts of Li, Na, K, Ca and Mg) or can form an organic salt (e.g., salts of triethyl amine, methylamine and ammonia),

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$$R_{17}$$
 R_{18} R_{20} R_{21} R

in which $R_{\rm B}$ and $R_{\rm B}$, which may be the same or different, each represent a hydrogen atom, an aliphatic group, an aromatic group, or a heterocyclic group that has the same meaning as defined for $R_{\rm B}$, or $R_{\rm B}$ and $R_{\rm B}$ may bond together to form a 5-to 7-membered ring; $R_{\rm B}$, $R_{\rm B}$, and $R_{\rm B}$, which may be the same or different, each represent a hydrogen atom, an aliphatic group, an aromatic group, or a heterocyclic group that has the same meaning as denied for $R_{\rm B}$, or an acyl group, an alkoxycarbonyl group, a sulfonyl group, a ureido group, or a urethane group, provided that at least one of $R_{\rm B}$ and $R_{\rm B}$ and at least one of $R_{\rm B}$ and $R_{\rm B}$ may bond together to form a 5-to 7-membered ring; at least two of $R_{\rm B}$, $R_{\rm B}$ and $R_{\rm B}$ may bond together to form a 5-to 7-membered ring;

R₁₀, R₁₁, R₁₂, R₁₃ and R₁₄, which may be the same or different, each represent a hydrogen atom, an aliphatic group (preferably having 2~74 carbon atoms, more preferably having 12~54 carbon atoms, e.g., methyl, isopropyl, t-butyl, vinyl, benzyl, octadecyl, and cyclohexyl), an aromatic group (preferably having 6~76 carbon atoms, more preferably having 12~56 carbon atoms, e.g., phenyl, pyridyl and naphthyl), a heterocyclic group (preferably having 2~60 carbon atoms, more preferably having 10~50 carbon atoms, e.g., piperidyl, pyranyl, furanyl and chromanyl), a halogen atom (e.g., chlorine and bromine), -SR₆, -OR₆ or - N - R₆

l P

(preferably having 1~60 carbon atoms), an acyl group (preferably having 2~60 carbon atoms, e.g., acetyl and benzoyl), an alkoxycarbonyl group (e.g., methoxycarbonyl, butoxycarbonyl, cyclohexylcarbonyl, and octyloxycarbonyl), an aryloxycarbonyl group (e.g., phenyloxycarbonyl and naphthyloxycarbonyl), a sulfonyl group (e.g., methanesulfonamido, and benzenesulfonamido), a sulfamoyl group, a ureido group, a urethane group, a carbamoyl group, a sulfo group, a carboxyl group, a nitro group, a cyano group, an alkoxalyl (e.g., methoxalyl, Isobutoxalyl, octyloxyxalyl, and benzoyloxyxalyl group), an aryloxalyl group (e.g., phenoxyxalyl and naphthoxyxalyl), a sulfonyloxy group (e.g., methanesulfonyloxy),

-P(R₈)₃, -P (R₈)₂, -P (R₈)₂, -P(OR₈)₃or a formyl group, wherein R₈ and R₉ each represent a hydrogen atom, an aliphatic group, an alkoxy group, or an aromatic group. The alkoxycarbonyl, aryloxycarbonyl, sulfonyl, sulfonamido,sulfamoyl, urethane group,carbamoyol, alkoxalyl, aryloxalyl group or sulfonyloxy group preferably has 1~60 carbon atoms. Of these compounds, those wherein the total of the Hammet sigma values is 0.5 or over for the group -SO₂M' are preferable in view of the effect of the invention.

Typical examples of the compounds represented by general formula (III) are given below:

(III-1)

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(-- - **)**

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(M-2)

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(II-3)

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NH₂—NHCOCH₃

OC₁₆H₃₃

(III-4)

30

SO₂HN(C₂H₅)

Cl

COC₁₅H₃₁(n)

35

(III-5)

40

© C₁₅H₃₁C NO₂

45

(M-6)

50

 $SO_2 \cdot \frac{1}{2} Ca^{2+}$ $O=P(OC_8H_{17}^{(n)})_2$

10 (III-7)

SO₂ Na $\begin{array}{c}
SO_2 \text{Na} \\
\hline
SO_2 C_{18} H_{37} \\
\end{array}$

(H-8)

SO₂Na SO₂C₁₈H₃₇ (w)

30

(III-9)

 SO_2N_2 $(^{(m)}C_6H_{13})_2P P(C_6H_{13}^{(m)})_2$

(III-10)

SO₂Na
CH₃

(M-11)

(III-12)

(M-13)

$$^{\text{(n)}}C_{15}H_{31}$$
 \sim SO₂Na

(III-14)

$$^{(n)}C_8H_{17}OC ^{(n)}C_8H_{17}OC ^{(n)}C_8H_{17}OC-$$

25

(Ⅲ-17)

(Ⅲ-18)

HS
$$C(CH_3)_4$$

NaS
$$\leftarrow$$
 C_8H_{17} (t)

45 (M-20)

HS
$$C_{12}H_{25}$$

(III-21)

5

10

25

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HO-C-NHCOCHO-C5H₁₁(t)
$$C_{2}H_{5}$$

$$C_{5}H_{11}(t)$$

20 (111-22)

$$HO-NHC- \bigcirc OC_{12}H_{25}$$

30 (III−23)

$$\begin{array}{c|c} C_2H_5 & OH \\ \hline C_2H_5 & CH \\ \hline CN \\ \hline CSH_{11} & CN \\ \hline CSH_{11}(t)O \\ \end{array}$$

(Ⅲ-24)

NaN₃

45 (III – 25)
$$OC_4H_9$$
 $C_8H_{17}(t)$

$$C_8H_{17}(t) OC_4H_9$$

(1 - 26)

$$C_{2}H_{5}$$

$$C_{4}H_{9}-CHCH_{2}NHC$$

$$C_{4}C_{4}H_{9}$$

$$C_{4}C_{4}CH_{2}CHC_{4}CH_{2}CHC_{4}CH_{9}$$

 $\begin{array}{c} \text{CH}_{3}-\text{CH}-\text{OC}-\text{C(CH}_{3})_{3} \\ \text{CONHCH}_{2}\text{CONHCH}_{2}\text{CH}-\text{C}_{4}\text{He} \\ \end{array}$

(II-28)

(**I**-29)

$$(CH_3)_3C-C-CH-O-C-O-C_2H_5$$
 SO_2
 $C_{10}H_{21}-NHOC$
 $CONH-C_{10}H_{21}$

(M-30)

5

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$$(t)C_{5}H_{11} - C_{5}H_{11}(t) - C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t) - C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t) - C_{5}H_{11}(t)$$

²⁰ (Ⅱ−31)

$$SO_2 \sim N^{\oplus}(C_4H_9(n))_4$$

$$C_{12}H_{25}-O-(CH_2)_3-NH-C$$

$$CNH(CH_2)_3O-C_{12}H_{25}(n)$$

$$O$$

(Ⅲ-32)

(**I**I-33)

55

(**1**-34)

5

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(□−35)

SO₂Li

$$C_5H_{11}(t)$$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$

(**I**-36)

35
$$SO_{2}NHN = H$$

$$CONH(CH_{2})_{3}O + C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t)$$

(II-37)

$$\begin{array}{c} \text{SO}_2\text{NHN} = \begin{pmatrix} H \\ H \end{pmatrix} \\ \text{C}_{12}\text{H}_{25}\text{OOC} \\ \end{array}$$

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(**I**I-38)

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(**I**-39)

SO₂NHN=
$$H$$

$$C_2H_5$$

$$C_2H_5$$

$$C_2H_5$$

$$C_2H_5$$

$$CONHCH_2CH-C_4H_9(n)$$

(**I**-40)

$$(t)C_{5}H_{11} - (C_{5}H_{11}(t)) - (C_{5}H_{11}($$

45 (II-41)

$$C_{16}H_{33}-O-C$$
 $C_{16}H_{33}$
 $C_{16}H_{33}$

5
$$C_{14}H_{29}-O-C$$
 $COOC_{14}H_{29}(n)$

CH3
$$SO_2N-NHSO_2-$$

$$C_2H_5$$

$$CONHCH_2CHC_4H_9(n)$$

$$C_2H_5$$

$$C_2H_5$$

$$C_2H_5$$

$$C_2H_5$$

$$\begin{array}{c} \text{OCH}_{3} \\ \text{C=O} \\ \text{SO}_{2}-\text{NH-N-SO}_{2}-\text{OOC}_{8}\text{H}_{17}\text{(n)} \\ \\ \text{OOC}_{8}\text{H}_{17}-\text{O-C} \\ \text{OOC}_{8}\text{H}_{17}\text{(n)} \end{array}$$

Synthesis Example 6 (Synthesis of exemplified compound III-30)

i) Synthesis of 3,5-di-(2,4-di-tertiary-acylphenoxypropylcarbamoyl)-benzenesulfonyl chloride

100 mt of toluene, 16 mt (0.080 mol) of a methanol solution containing 28 % of sodium methylate and 24.7 g (0.085 mol) of 2,4-di-tertiary-amylphenoxypropylamine were added to 10 g (0.034 mol) of sodium dimethyl 5-sulfoisophthalate, followed by heating to 100°C. The reaction mixture was heated for 3 hours while the methanol was distilled off, and after cooling, cold water was added thereto. The separated toluene layer was washed twice with cold water and then dried over Glauber's salt. The Glauber's salt was filtered out, followed by condensation, and to the resulting dried concentrate were added 100 mt of N,N-dimethylacetamide (DMAC) and 50 mt of acetonitrile to dissolve it, followed by stirring at room temperature. 30 mt (0.326 mol) of phosphorus oxychloride was added thereto, followed by heating to 50 to 60°C for 1 hour. Ice-water was added to the reaction mixture, extraction was carried out with 300 mt of ethyl acetate, and the ethyl acetate layer was washed three times with cold water and then dried over Glauber's salt. The Glauber's salt was filtered out, the ethyl acetate was removed by distillation, and the product was purified by column chromatography. The yield was 11.5 g (41.9 % of theory).

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ii) Synthesis of sodium 3,5-di-(2,4-ditertiaryaminopheoxypropylcarbaboyl)-benzenesulfinate (exemplified compound III-30)

100 mt of water and 20 mt of acetonirile were added to 2 g (0.016 mol) of sodium sulfite and 2.4 g (0.029 mol) of sodium hydrogen carbonate, followed by stirring at 30°C. A solution of 10.5 g (0.013 mol) of 3,5-di-(2,4-ditertiary-amylphenoxypropylcarbamoyl)-benzenesulfonylchloride in 100 mt of acetonitrile was added thereto dropwise. After stirring for 1 hour it was poured into ice-water, followed by extraction with 150 mt of ethyl acetate. The ethyl acetate layer was washed with cold water tree times and then dried over Glauber's salt. The Glauber's salt was filtered out, followed by concentrating to dryness to produce 8.6 g of exemplified compound III-30 as a solid in a yield of 82.8 %. Results of elementary analysis (C₄₆H₆₇N₂O₆SNa)

Hesuits of elementary analysis (C46H6N2O6SNa)

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15		С	Н	N	S
	Found (%)	68.75	8.39	3.32	3.92
	Calculated (%)	69.14	8.45	3.51	4.01

Synthesis Example 7 (Synthesis of exemplified compound III-41)

15 mt of ethyl acetate was added to 1.0 g of 3,5-dihexadecyloxycarbonylbenzenesulfonyl hydrazide and 5 mt of dimethylacetamide, followed by stirring at room temperature, and 1.01 g of crystals of 3,5-dihexadecyloxycarbonylbenzenesulfonic acid chloride was added thereto. After stirring for 30 min at room temperature 0.2 mt of pyridine was added thereto, followed by stirring for 5 hours. After the completion of the reaction the reaction liqid was poured into 100 mt of water, and the deposited crystals were filtered and dried. The crystals were purified by silica gel column chromatography to produce 0.4 g (20.5 %) of crystals of exemplified compound III-41, melting point 148 to 150°C.

The reaction between the aromatic amine developing agent and the compound represented by general formula (I) or (II) can be shown by the following formula (a), and the reaction between the oxidized product of the aromatic amine developing agent and the compound represented by general formula (III) can be shown by the following formula (2). These reactions proceed gradually during the storage of a color photograph.

(1)
$$R_{1}-(A)_{\overline{n}}-X + NH_{2}-(R_{24}) \longrightarrow R_{1}-(A)_{\overline{n}}-NH-(R_{24}) \longrightarrow (R_{23})_{5}$$

$$R_{2}-C=Y + NH_{2} \xrightarrow{R_{24}} -R_{24} \rightarrow R_{2}-C-YH$$

$$R_{23})_{s}$$

$$(I)$$

50

$$R_2-C=N-\begin{array}{|c|} & & & \\ & \downarrow & \\ & &$$

or

$$\begin{array}{c}
Z - R_7 \\
NH_2 \longrightarrow R_2 \\
R_{23} \\
R_{23} \\
\end{array}$$

In formulae (1) and (2) shown above, the groups in general formulae (I) to (III) have the same meaning as defined above. R₂₁ represents a hydrogen atom, an alkyl group (including a substituted alkyl group, e.g., methyl, ethyl or hydroxymethyl), or an alkoxy group (including a substituted alkoxy group, e.g., methoxy, ethoxy, or methoxyethoxy); R₂₄ represents a hydroxy group or an amino group (including a substituted amino group, e.g., amino, N-methylamino, N,N-dimethylamino, N,N-diethyl, N-ethyl-N-(2-methanesulfonamidoethyl)amino, N-ethyl-(2-hydroxyethyl)amino and N-ethyl-N-(2-methoxyethyl)amino); and s is an integer of 1 to 4.

In the present invention, if compound (A) or (B) has a low molecular weight or is readily soluble in water, it may be added to a processing solution so that the compound may be taken into the photographic material during the development processing. It is a preferable method wherein the compound is added to the hydrophilic colloid layer of the photographic material in a step of the production of the photographic material. In the latter method, the compound is dissolved in a single high-boiling solvent (oil) that has a boiling point of 170°C or over at atmospheric pressure, or a single low boiling solvent, or a solvent mixture of said oil and a low boiling solvent, and the resulting solution is emulsified and dispersed in an aqueous hydrophilic colloid solution, such as an aqueous gelatin solution. In the present invention it is preferable that compound (A) or (B) is dissolvable in a high boiling organic solvent. The particle diameter of this emulsified dispersion is not particularly limited, but preferably the particle diameter is 0.05 to 0.5 μm, more preferable

0.1 to 0.3 µm. It is preferable that compound (A) or (B) be co-emulsified with a coupler. In this case the oil/coupler weight ratio is preferably from 0.01 to 2.0.

In the present invention the proportion of compound (A) or (B) is such that 1×10^2 to 10 mol, preferably 3×10^2 to 5 mol, be present per mol of a coupler. If the amount of compound (A) or (B) is too small, the exhibited effect of the invention tends to lower, whereas if the amount of compound (A) or (B) is too large, the color forming reaction is liable to be hampered or the decomposition of compound (A) or (B) becomes noticeable and tends to damage the color image. In particular, the amount of compound (B) added is preferably in the range of 2×10^2 to 2×10^1 per mol of a coupler.

Specific examples of the above-mentioned oils include alkyl phthalates (e.g., dibutyl phthalate, dioctyl phthalate, diisodecyl phthalate, and a dimethoxyethyl phthalate), phosphates (e.g., diphenyl phosphate, triphenyl phosphate, triphenyl phosphate, tricresyl phosphate, dioctylbutyl phosphate, and monophenyl-p-t-butylphenyl phosphate), citrates (e.g., tributyl acetylcitrate), benzoates (e.g., octyl benzoate), alkylamides- (e.g., diethyl-laurylamide and dibutyllaurylamide), fatty acid esters (e.g., dibutoxyethyl succinate, diethyl azelate, and dioctyl sebasate), trimesate (e.g., tributyl trimesate), compounds containing epoxy rings (e.g., compounds described in U.S. Patent No. 4,540,657), and phenols (e.g.,

$$HO - C_5H_{11}(t)$$
 , $HO - C_{12}H_{25}(t)$, $C_{12}H_{25}(t)$

$$C_5H_{11}(t)$$
 $C_5H_{11}(t)$

HO
$$C_8H_{17}(t)$$
 $C_8H_{17}(n)$
 $C_8H_{17}(n)$
 $C_8H_{17}(n)$
 $C_8H_{17}(n)$
 $C_8H_{17}(n)$
 $C_8H_{17}(n)$
 $C_8H_{17}(n)$

ethers (e.g., phenoxyethanol, and diethylene glycol monophenyl ether). Low boiling solvents used as auxiliary solvents include organic solvents that have a boiling point of about 30°C-150°C under atmospheric pressure, such as lower alkyl acetates (for example, ethyl acetate, isopropyl acetate, and butyl acetate)

ethyl propionate, methanol, ethanol, sec-butyl alcohol, cyclohexanol, fluorinated alcohols, methyl isobutyl ketone, β -ethoxyethyl acetate, methyl cellosolve acetate acetone, methyl acetone, acetonitrile, dioxane, dimethylformamide, dimethylsulfoxide, chloroform, and cyclohexane.

Instead of high boiling organic solvents, not only oil solvents (including ones that are solid at room temperature, such as waxes) that are additives for couplers, etc., but also latex polymers can be used, and additives such as couplers, color mixing preventive agents, and ultraviolet absorbing agents can also serve as oil solvents.

As latex polymers, use can be made of latex polymers produced by using one or more monomers such as acrylic acid and methacrylic acid, and their esters (e.g., methyl acrylate, ethyl acrylate and butyl acrylate), acrylamide, t-butylacrylamide, methacrylamide, vinyl esters (e.g., vinyl acetate, and vinyl propionate), acrylonirile, styrene, divinylbenzene, vinyl alkyl ether (e.g., vinyl ethyl ether), maleic acid esters (e.g., methyl maleate), N-vinyl-2-pyrrolidone, and N-vinyl pyridine, 2-and 4-vinyl pyridine.

In the present invention, examples of surface active agents used for dispersing into an aqueous protective colloid solution, solutions in which compound (A) or (B) optionally with a coupler is dissolved include saponin, sodium alkylbenzenesulfosuccinates, and sodium alkylbenezenesulfonates.

Preferably anionic surface active agents having a sulfonic acid group such as compounds shown below are used alone or in combination:

Preferred combination of compounds (A) and (B) for use in the present invention is the combination of a compound (A) selected from compounds represented by general formula (I) and a compound (B) selected from compounds represented by general formula (IV), and especially preferred combination is that of a compound (A) selected from compounds represented (I-a) or (I-c) and a compound (B) selected from compounds represented by general formula (IV). A combination of compounds (A) and (B) selected from compounds represented by general formula (I-a) and (IV) respectively is most preferable.

In the present invention, compound (A) or (B) may be added to any of a color developing solution, a bleaching solution, a fixing solution, a washing solution, and a rinsing solution. In this case, the concentration of compound (A) or (B) in the processing solution is 10 ⁵ mol/1 to 10 ¹ mol/1.

The compound of the present invention can be used together with the following oxidation inhibitors and fading preventive agents.

Representative patents in which these fading preventive agents and oxidation inhibitors are described include: U.S.Patent Nos. 3,935,016, 3,982,944, 3,700,455, 3,764,337, 3,432,300, 3,574,627, 3,573,050, and 4,254,216, Japanese Patent Application (OPI) Nos. 21004/1980, 145530/1979, 152225/1977, 20327/1978, 17729/1978, 72246/1986, 73152/1986, 90155/1986, 90156/1986, 145554/1986, and 6321/1980, Japanese Patent Publication Nos. 12337/1979, and 31625/1973, British Patent No. 1,347,556, and British Patent Application (OPI) No. 2,066,975.

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 $\begin{array}{c|c} CH_3 \\ CH_3 \\ C_3H_7 - C \end{array} \begin{array}{c} CH_3 \\ CH_3 \\ CH_3 \end{array}$

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CH₃ CH₂ CH₂ CH₂ CH₁₃ COOC₆H₁₃ CH₃ CH₃ OH

CH₃
CH₃
CH₃
CH₃
CH₃
CH₃
CH₃
CH₃

С́Н₃ ОСН₃

OCH₃ CH₃
CH₃ CC-C₃H₇
CH₃ OCH₃

CH₃ CH₃

CH₃ CH₃

CH

CH

CH

CH₃

CH₃

CH₃

CH₃ CH₃ CH₃

CH₃ CH₃ CH₃

CH₃ CH₃ CH₃ C₈H₁₇(t)

(t)C₈H₁₇ CH₃ CH₃

CH₃ CH₃ CH₃ CH₃ CH₃ OCH₃

$$CH_{2} CH_{2} CH_{2}O - (CH_{2})_{2} - O - CH_{2} CH_{2}O$$

CH₃ OH OH CH₃

$$CH_3 - C + CH_2 + CH_3$$

$$CH_3 - CH_3 + CH_3$$

$$CH_3 - CH_3 + CH_3$$

 $\begin{bmatrix}
C_2H_5 & C_2H_5 \\
C_4H_9 - CHCH_2O & OCH_2CH - C_4H_5 \\
C_2H_5 & C_2H_5 \\
OCH_2CH - C_4H_5 \\
C=N & Ni \\
C=N & OH
\end{bmatrix}$

$$\begin{array}{c}
NH_{2}C_{12}\dot{H}_{25} \\
\downarrow \\
O - Ni - O \\
\hat{S} \\
\downarrow \\
C_{8}H_{17}(t) C_{8}H_{17}(t)
\end{array}$$

$$(t)C_8H_{17} \longrightarrow O \qquad HO \longrightarrow C_8H_{17}(t)$$

$$S \rightarrow Ni \leftarrow S$$

$$(t)C_8H_{17} \longrightarrow OH \qquad O \longrightarrow C_8H_{17}(t)$$

$$C_{4}H_{9} \xrightarrow{CH_{3}} C_{4}H_{9} \xrightarrow{CH_{3}} C_{4}H_{9}$$

$$C_{4}H_{9} \xrightarrow{CH_{3}} C_{4}H_{9}$$

$$C_{4}H_{9} \xrightarrow{CH_{3}} C_{4}H_{9}$$

 $C_4H_9(t)$ $C_4H_9(t)$ $C_4H_9(t)$ $C_4H_9(t)$

 $HO - CH_2CH_2CO_2C_{18}H_{17}$ C_4H_9

CH₃ CH₃
HN -OC-C₇H₁₅
CH₃ CH₃ 0

$$\begin{pmatrix}
C_4H_9(t) \\
HO \longrightarrow CH_2 \\
C_4H_9(t)
\end{pmatrix}_2 C \leftarrow \begin{pmatrix}
CH_3 & CH_3 \\
CO_2 \longrightarrow N-COCH_3 \\
CH_3 & CH_3
\end{pmatrix}_2$$

$$\left(\begin{array}{c}
\text{HO} - \left(\begin{array}{c}
\text{CH}_3 \\
\text{CH}_3
\end{array}\right) \\
\text{CH}_3 \\
\text{CH}_3
\end{array}\right)_2$$

$$N(C_4H_9)_2$$

$$C_4H_9O \longrightarrow C_8H_{17}(t$$

CH₃

$$C_{16}H_{33} - N O$$

$$CH_2$$

$$C_8H_{17}$$
 $C_8H_{17}O$
 $C_4H_9(t)$

 $C_{14}H_{29}-N$

 $C_{14}H_{29}-N$ $N-C_{14}H_{2}$

$$C_{12}H_{25}-N$$

Ultraviolet absorbers that can be used in the present invention include those listed in Research Disclosure (R.D.) No. 17643, VII-C, and preferably are benzotriazole derivatives represented by the following general formula (XVII):

General formula (XVII)

In the formula R_{51} , R_{52} , R_{53} , R_{54} , and R_{65} , which may be the same or different, each represent a hydrogen atom, an alkoxy group, an alkyl group, a halogen atom, or an alkoxycarbonyl group.

$$\bigcap_{N} \bigvee_{N \to C_4H_9(t)} \bigvee_{N$$

Al

55 .

 $C\ell \bigvee_{N} \bigvee_{N} C_4H_9(t)$

$$\begin{array}{c|c}
 & OH \\
 & C_4H_9(t) \\
 & CH_2CH_2COOC_6H_{13}
\end{array}$$

The process for the production of a color photograph of the present invention is the same as the conventional process for the production of a color photograph, except that compound (A) or (B) is employed in the manner described above.

Color photographic materials to which the present process for the production of a color photograph will be applied are not particularly limited, and typical examples of the color photographic materials include color papers, color negative film for general purposes and movies, color reversal films for slides and television, color positive films, and color reversal papers. The present invention can also be applied to black

and white photographic materials that use a mixture of three color couplers, as described in Research Disclosure 17123 (June 1978).

Therefore, there is no particular limit to the couplers employed in the color photographic materials, and examples include:

(a) Yellow couplers

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Couplers represented by the general formulae (Y-I) and (Y-II):

General formula (Y-I)

С Н ₃ - С - С О С Н - R ₁₁

wherein R₁₁ represents a substituted or unsubstituted N-phenylcarbamoyl group, and Z₁₁ represents a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent.

General formula (Y-II)

$$(R_{12})_s \qquad \begin{matrix} COCH-R_{11} \\ Z_{11} \end{matrix}$$

wherein $R_{\rm H}$ represents a substituted or unsubstituted N-phenylcarbamoyl group, $Z_{\rm H}$ represents a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent, $R_{\rm H}$ represents a hydrogen atom or a substituent group, and s is an integer of 1 to 5.

In more detail, typical chemical structures of the yellow couplers represented by general formulae (Y-I) and (Y-II) are the same ones as described, for example, in U.S. Patent Specifications given below, wherein the numbers in parentheses indicate the columns describing the chemical structures: U.S.Patent Nos. 3,894,875 (1-2), 3,408,194 (2-3), 4,404,274(3-17), 4,022,620 (3-7) and 4,057,432 (1-4).

(b) Magenta couplers

Couplers represented by the following general formulae (M-I) and (M-II):

General formula (M-I)

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wherein R₂₁ represents an alkyl group, an aryl group, or a carbamoyl group; Ar represents a phenyl group or a phenyl group substituted by one or more of halogen atoms, alkyl groups, cyano groups, alkoxy groups, alkoxycarbonyl groups, or acylamino groups; and Z₂₁ represents a hydrogen atom or a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent.

General formula (M-II)

$$\begin{array}{c|c} R_{22} & Z_{21} \\ \hline \\ N & Z_{22} \\ \hline \\ Z_{22} & Z_{2} \end{array}$$

wherein R_{22} represents a hydrogen atom or a substituent group; Z_{21} represents a hydrogen atom or a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent; Z_{22} , R_{22}

 Z_{23} and R_{24} represent - C = -N = or -NH-; one of the $Z_{27}Z_{23}$ bond and the $Z_{27}Z_{22}$ bond is a double bond and the other is a single bond; and when the $Z_{27}Z_{22}$ is a carbon-carbon double bond, the double bond may be part of an aromatic ring.

In more detail, typical chemical structures of the magenta couplers represented by general formulae (M-I) and (M-II) are the same ones as described in U.S.Patent Specifications, etc. given below, wherein the numbers in parentheses indicate the columns or the pages describing the chemical structures: U.S.Patent Nos. 3,519,429 (2-6), 3,558,319 (2-3), 3,725,067 (2-8), 3,935,015 (3-7), 4,241,168(2-14), 4,351,897 (2-6), 4,367,282 (3-10), and 4,540,654 (2-8), Japanese Patent Application (OPI) No. 65245/1986 (pages 378-384), and WO-86-1915 (pages 5-10).

(c) Cyan couplers

Cyan couplers represented by the general formula (C-I):

General formula (C-I)

$$R_{33} \xrightarrow{OH} NHCOR_{31}$$

$$R_{32} \xrightarrow{Z_{31}}$$

wherein R₃₁ represents an alkyl group, a cycloalkyl group, an aryl group, an amino group, or a heterocyclic group; R₃₂ represents an acylamino group or an alkyl group; R₃₃ represents a hydrogen atom, an alkyl group or an alkoxy group; R₃₃ and R₃₂ may bond together to form a ring; and Z₃₁ represents a hydrogen atom, a halogen atom, or a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent.

In more detail, typical chemical structures of the cyan couplers represented by general formula (C-I) are the same ones as described, for example, in U.S.Patent Specifications giben below, wherein the numbers in parentheses indicates the columns describing the chemical structures: U.S.Patent Nos. 2,920,961 (1), 3,772,002 (1-3), 3,864,366 (2-6), 4,124,396 (2), 4,333,996 (2-8), 4,565,777 (3-5), and 4,564,586 (2-4).

The above couplers may form a dimer or an even higher polymer.

Preferred couplers for use in combination with preservability improving compounds (A) and (B) of the present invention are couplers represented by general formula (Y-I), (M-II) or (C-I), of which especially preferable being couplers represented by general formula (M-II) or (C-I).

Specific examples of these couplers are given below, but the invention is not limited to them.

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(Y-1) $C_5H_{11}(t)$ $NHCOCHO - C_5H_{11}(t)$ C_2H_5 $O - C_{13}$ CH_3

(Y-2)

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$$(CH_3)_3C-COCHCONH-CI
OVNOCHCOC12H25(n)
N-CH-OC2H5$$

(Y-3)

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$$(CH_3)_3C-CO CHCONH$$

NHCO $(CH_2)_3O$
 $C_5H_{11}(t)$

SO₂

OCH₂

OCH

$$(Y-4)$$

$$(CH_3)_3C-COCHCONH$$

$$O$$

$$NHCO(CH_2)_3O$$

$$C_5H_{11}(t)$$

$$COOH$$

(Y-5)

$$(CH_3)_3C-COCHCONH \longrightarrow C_5H_{11}(t)$$

$$O \searrow N \searrow O$$

$$N-CH_2$$

$$N+CH_2$$

$$O \searrow N \longrightarrow C_5H_{11}(t)$$

$$N+CH_2$$

$$(Y-6)$$

$$(CH_3)_3C-COCHCONH-C_1$$

$$C_4H_9$$

$$COOCHCOOC_{12}H_{25}$$

$$CH_2-C_1$$

(CH₃)₃C-CO CHCONH-COC₁₆H₃

$$(CH_3)_3 C - CO CHCONH - COCC16H3$$

$$(CH_3)_3C-COCHCONH- NHCOC(CH_3)_3$$

$$CN O-CH_2CH_2-O-C_5H_{11}(t)$$

$$COOCH_3$$

CL

$$C_5H_{11}(t)$$

 $C_5H_{11}(t)$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$
 $C_5H_{11}(t)$

$$(Y-15)$$

(Y-14)

$$(CH_3)_3C-COCHCONH$$

$$O \downarrow N \downarrow O$$

$$N-CH-OC_{16}H_{33}$$
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$$(Y-16)$$

$$(CH_3)_3C-COCHCONH-C$$

$$NHCO(CH_2)_3O-C_5H_{11}(t)$$

$$NHCO(CH_2)_3O-C_5H_{11}(t)$$

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$$\begin{array}{c} \text{CH}_{3}\text{O} & \text{CY-17}) \\ \text{CH}_{3}\text{O} & \text{COCHCONH-} \\ \text{O} & \text{N} & \text{O} \\ \text{CH}_{3} & \text{COCHCONH-} \\ \text{CH}_{3} & \text{CH}_{3} & \text{CH}_{3} \\ \text{CH}_{4} & \text{CH}_{3} & \text{CH}_{4} \\ \text{CH}_{4} & \text{CH}_{4} & \text{CH}_{4} & \text{CH}_{4} \\ \text{CH}_{4} & \text{CH}_{4} & \text{CH}_{4} \\ \text{CH}_{4} & \text{CH}_{4} & \text{CH}_{4} \\ \text{CH}_{4} & \text{CH}$$

$$(Y-18)$$

$$Cl$$

$$CH_3O- \bigcirc COCH CONH - \bigcirc C_2H_5$$

$$O \bigcirc N \bigcirc O$$

$$HC-N$$

$$C_2H_5O$$

$$CH_2- \bigcirc C_5H_{11}(t)$$

$$C_5H_{11}(t)$$

$$(Y-19)$$

$$CH_{3}O- (Y-19)$$

$$CD-CHCONH- (Y-19)$$

$$O-CHCONH- (Y-19)$$

$$O-$$

$$\begin{array}{c|c} C_{16}H_{33}O \longrightarrow & COCHCONH \longrightarrow & SO_2N \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\$$

(Y-21)

(Y-22)

$$CH_3O \longrightarrow COCHCONH \longrightarrow COOC_{12}H_{25}$$

$$O \longrightarrow N \longrightarrow O$$

$$COOC_{12}H_{25}$$

$$O \longrightarrow N \longrightarrow CH$$

$$CH_3 \longrightarrow OC_6H_{13}$$

CL
CH₃O-
$$\langle - \rangle$$
-COCHCONH- $\langle - \rangle$
NHCO COOCH₃
 $\langle - \rangle$ -CH₂ \rangle -CH₂- $\langle - \rangle$ -CH₂ \rangle -CH₃

$$CH_3O - COCHCONH - COOC_{12}H_{25}$$

$$O + O + COOC_{12}H_{25}$$

$$CH_2 - COOC_{12}H_{25}$$

(Y-25)

$$(CH_3)_3C - COCHCONH - C_4H_9$$

$$COOCHCOOC_{12}H_{25}$$

$$CH_2 - COCHCOOC_{12}H_{25}$$

(Y-26)

$$(CH_3)_3C - COCHCONH - CH_3$$

$$O \downarrow N \downarrow O$$

$$NHCOCHCH_2 SO_2 C_{12}H_{25}$$

$$CH_3$$

$$NHCOCHCH_2 SO_2 C_{12}H_{25}$$

(Y-2.7)

$$(CH_3)_3C - COCHCONH - COO(CH_2)_2 OC - C_4H_9(t)$$

$$O = OC_2H_5$$

$$OC_2H_5$$

$$OC_2H_5$$

(Y-28)

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$$(t)C_{5}H_{11} \xrightarrow{} O - CH_{2}CNH$$

$$C_{5}H_{11}(t) O CL$$

$$C_{15}C$$

(M-2)20

$$C_{2}H_{5} \longrightarrow CONH$$

$$C_{5}H_{11} \longrightarrow C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t)$$

$$C_{6}U$$

30 (M-3)

$$(M-3)$$

$$C_{2}H_{5}$$

$$C_{15}H_{31}$$

$$CH_{3}O$$

$$CH_{3}$$

$$CH_{3}$$

(t)
$$C_5H_{11}$$
 — O-CHCONH NO CL OCH₃

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(M-7)

 $C_{13}H_{27}CONH$ N Cl Cl

(M-8)

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$$(M-9)$$

$$Cl$$

$$C_{12}H_{25}$$

$$NH$$

$$N$$

$$Cl$$

$$Cl$$

$$Cl$$

$$Cl$$

$$Cl$$

$$Cl$$

$$Cl$$

$$(M-10)$$

³⁰ (M-11)

35 (t)
$$C_5H_{11}$$
 $O-(CH_2)_3-O-C$ NH O CL CL CL

(M-12)

$$(M-13)$$

$$(t) C_{5}H_{11} - C_{5}H_{11}(t) - C_{5}H_{11}$$

$$(M-17)$$

$$(M-18)$$

$$(M-19)$$

CL
$$C_{41}$$
 C_{45} C_{13} C_{13}

$$(M-20)$$

$$Cl \qquad OC_4H_9$$

$$Cl \qquad NH \qquad S-Cl$$

$$Cl \qquad CR$$

$$Cl \qquad NH \qquad S-Cl$$

$$Cl \qquad CR$$

$$Cl \qquad COOC_2H_5$$

$$S-CH-C_{12}H_{25}$$

$$Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl \qquad Cl$$

$$Cl \qquad Cl \qquad Cl \qquad Cl \qquad Cl$$

$$(M-24)$$

$$C_{16}H_{33} - N - C_{2}H_{5}$$

$$(M-25)$$

$$C_{16}H_{33} - N - C_{2}H_{5}$$

$$C_{16}H_{33} - N - C_{2}H_{5}$$

$$C_{16}H_{33} - N - C_{2}H_{5}$$

$$C_{16}H_{33} - C_{16}H_{17}$$

$$N_{10} - C_{16}H_{17$$

$$(M-28)$$

$$CH_{3}$$

$$NH$$

$$CCP$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{3}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{3}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{3}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{3}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

$$CH_{4}$$

$$CHCH_{2}NHSO_{2}$$

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C₈H₁₇(t)

(M-32)

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$$CH_3 \longrightarrow C\ell$$

$$N \longrightarrow NH$$

$$C_4H_9$$

$$CH_2)_2NHC-CHO \longrightarrow C_5H_{11}(t)$$

$$C_5H_{11}(t)$$

(M-33)

$$HO - \left\langle \begin{array}{c} C_{10}H_{21} \\ \\ O - CH CNH - \left\langle \begin{array}{c} CH_2 \\ \\ O \end{array} \right\rangle \\ O - CH_2 \\ O - CH$$

(M-34)

(M - 35)

50
$$CH_{3} C\ell$$

$$N' NH$$

$$N NH$$

$$CH_{3} C\ell$$

$$N' NH$$

$$CH_{3} C\ell$$

$$N' NH$$

$$CH_{21} CH_{21} C\ell$$

$$N' NH$$

(M-36)

$$\begin{array}{c} \text{CH}_{3} \\ \text{CH}_{3} \\ \text{CH}_{3} \\ \text{CH}_{3} \\ \text{OC}_{4}\text{H}_{9} \\ \text{N} \\ \text{NH} \end{array}$$

$$(M-38)$$

$$(M-39)$$

$$\begin{array}{c}
OC_4H_9 \\
-SO_2NH - (CH_2)_3 \\
N \\
N \\
NH
\end{array}$$

(M-41)

$$(M-42)$$

$$OC_8H_{17}(n)$$

$$OC_8H_{17}(n)$$

$$OC_8H_{17}(n)$$

$$OC_8H_{17}(n)$$

$$(M-43)$$

$$OCH_3 OC_4H_9(n)$$

$$C_8H_{17}(t)$$

$$N= (CH_2)_2NHSO_2$$

$$C_8H_{17}(t)$$

$$C_8H_{17}(t)$$

₂₀ (M-44)

OCH₃ OC₄H₉(n)
$$C_8H_{17}(t)$$
NHSO₂

$$C_8H_{17}(t)$$

$$C_8H_{17}(t)$$

(C-3)

(C-4)

$$(t)C_4H_9 \xrightarrow{C_2H_5} NHCOCHO \xrightarrow{C_2H_5} (t)C_5H_{11}$$

OH
$$C_{2}H_{5}$$

$$OCH_{2}CH_{2}CH_{2}COOH$$

$$OCH_{2}CH_{2}CH_{2}COOH$$

$$(t)C_5H_{11} \longrightarrow OCHCONH$$

$$(t)C_5H_{11}$$

$$(t)C_5H_{11}$$

(C-8)
$$C_{12}H_{25}$$
OH
NHCO
(t)C₄H₉

$$C\ell$$

20 (C-9)

$$(t)C_5H_{11} - \underbrace{\begin{array}{c} C_6H_{13} \\ OCHCONH \\ C\ell \end{array}} NHCO - \underbrace{\begin{array}{c} C_\ell \\ C\ell \end{array}}$$

(C-11)

$$C_{6}H_{13}$$
OH
$$C_{6}H_{13}$$
NHCO
$$HNSO_{2}C_{4}H_{6}$$

(C-12)

$$O \longrightarrow C_8H_{17} \longrightarrow OCHCONH \longrightarrow HNSO_2CH_2CH_2OCH_3$$

$$(t)C_6H_{13}$$

20 (C-13)

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OH NHCO
$$\longrightarrow$$
 (t) C_5H_{11}
HNSO₂(CH₃)₄O \longrightarrow (t) C_5H_{11}

(C-14)

(C-15)

 $O = \begin{pmatrix} H & OH \\ N & NHCO - \\ N & HNSO_2 - \\ OCH_2CHC_4H_9 \\ C_2H_5 \end{pmatrix}$

²⁰ (C-17)

$$O = \bigvee_{\substack{N \\ H}} OH$$

$$O = \bigvee_{\substack{N \\ C2}} NHCO - \bigvee_{\substack{C_2 H_5 \\ NHCO \ CH \ O}} (t)C_5H_{11}$$

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(c-18)

 $O = \underbrace{\begin{array}{c} CH_3 \quad CH_3 \\ OH \\ NHCO - \\ NHSO_2C_{16}H_{33}(n) \end{array}}_{NHSO_2C_{16}H_{33}(n)}$

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(C-19)

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15 OH OH NHCONH
$$-CN$$

C₄H₉ NHCONH $-CN$

(t) C₅H₁₁ OCHCONH

20 (C-21)

$$(t)C_8H_{17} - (t)C_8H_{17}$$

$$(t)C_8H_{17} - (t)C_8H_{17}$$

$$(t)C_8H_{17}$$

$$(t)C_8H_{17}$$

$$(t)C_8H_{17}$$

(c-22)
$$C_2H_5 \qquad \text{OH}$$

$$C_2H_5 \qquad \text{NHCONH} \longrightarrow SO_2C_3H_5$$

$$(t)C_5H_{11} \longrightarrow OCHCONH$$

(c-23)
$$C_4H_9 \longrightarrow NHCONH \longrightarrow CN$$
(t) $C_5H_{11} \longrightarrow OCHCONH \longrightarrow CN$

$$(C-24)$$

$$C_{4}H_{9}$$

$$(t)C_{5}H_{11}$$

$$OCHCONH$$

$$OCH_{3}$$

$$(C-25)$$

$$(C-25)$$

$$(C-25)$$

$$(C-25)$$

$$(C-25)$$

$$(C-26)$$

$$(C-26$$

$$\begin{array}{c} \text{OH} & C_2H_5 \\ \text{Cl} & \text{NHCOCHO} & C_5H_{11}(t) \\ \text{CH}_3 & \text{Cl} & \text{Cl} & \text{Cl} \\ \end{array}$$

$$(C-28)$$

$$Cl$$
 Cl
 Cl
 Cl
 Cl
 Cl
 Cl

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(C-29)

OH
$$C_4H_9$$

$$C_5H_{11}$$

$$C_5H_{11}(t)$$
OH
$$NHCONH$$

$$CN$$

25 (C-30)

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(C-31)

OH

$$C\ell$$

NHCOCH₂O $C_5H_{11}(t)$

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The process for the production of a color photograph of the present invention is the same as the common process for the production of a color photograph, except that compound (A) or (B) is contained suitably in a processing solution.

The silver halide grains used in the present invention may be in the form of regular crystals, such as cubic crystals, octahedral crystals, dodecahedral crystals, and tetradecahedral crystals, or of irregular crystals, such as spherical crystals, or in a tabular form having a length/thickness ratio of 5 or more. The emulsion may comprise a composite of these crystalline forms or a mixture of them.

The composition of the silver halide comprises silver chloride, silver bromide, or a mixed silver halide, and the silver halide that is preferably used in the present invention is silver chloro(iodo)(bromide, silver (iodo)chloride or silver (iodo)bromide that contains no silver iodide, or a maximum of 3 mol % of silver iodide if it is contained.

The average grain size of the silver halide grains is preferably a maximum of 2 μ m and a minimum of 0.1 μ m, more preferably a maximum of 1.3 μ m and a minimum of 0.15 μ m. The grain size distribution may

be narrow or wide, although in the present invention it is preferable to use the "monodisperse" silver halide emulsion having narrow grain size distribution wherein 95% or over of all the grains fall within ±40%, preferably ±30%, and more preferably ±20% of the average grain size in terms of the number of grains or in terms of weight with a view to improving graininess and the sharpness. In order to satisfy the gradation at which the photographic material is aimed, in emulsion layers having substantially identical color sensitivity two or more monodisperse silver halide emulsions different in grain size, or grains having the same size but different in sensitivity, are mixed and applied in the same layer or are applied as separate layers. Further, a combination of two or more polydisperse silver halide emulsions or a combination of a monodisperse emulsion and a polydisperse emulsion can be mixed or applied as separate layers.

In the silver halide emulsion used in the present invention, the inside or the surface of the grains may be chemically sensitized, for example for sulfur sensitization, selenium sensitization, reduction sensitization, or noble metal sensitization, which may be used alone or in combination. Detailed examples thereof are described, for example, in patents cited in Research Disclosure No. 17643-II (Dec.1978), page 23.

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The silver halide emulsion used in the present invention can also be spectrally sensitized in a conventional manner using a cyanine dye or a merocyanine dye.

Gelatin to be used in this invention may be an alkaline-treated gelatin (having an isoelectric point of 4.5 to 5.3), an acid-treated gelatin (having an isoelectric point of 6.0 to 9.5) or an enzyme-treated gelatin. Of these, the acid-treated gelatin is preferable in view of the prevention of stain. Preffered amount of acid-processed gelatin in total coating amount of gelatin to be added is 10% or more, with being more preferrably 25% or more, and most preferrably 50% or more, and the upper limit is 100%.

It is preferable that the color developing solution used in the present invention be substantially free from benzyl alcohol. When a low-replenishing type color development replenishing solution is prepared, if benzyl alcohol is contained in some cases it takes a longer time to dissolve the components due to the slow dissolving rate, or a tarry substance is formed. On the other hand, even if a color developing solution free from benzyl alcohol is of a low-replenishing type, since the components can be dissolved within a short period of time and a tarry substance will not be formed, it is easy and advantageous to prepare a low-replenishing type development replenisher. When continuous processing is effected by using a color developing solution free from benzyl alcohol, which is prevented from the composition fluctuation of the solution the replenishing amount can be lowered to half or below (165 mt/m² or below) the standard replenishing amount, and constant finishing can be obtained without the formation of tarry substances or a change of stain.

As additives used in the color developing solution, use can be made of various compounds described in Japanese Patent Application Nos. 1667/1984, pages 14 to 22, 118418/1984, pages 45 to 50, and 32462/1986, pages 11 to 22.

As antifoggants to be used in the color developing solution, use can be made of tetrazaindenes, benzoindazoles, benzoindazoles

The photographic emulsion layer after the color development is generally subjected to a bleaching process. The bleaching process may be carried out as a one-bath bleach/fix simultaneously with a fixing process, or it may be carried out separately from the fixing process. In order to quicken the processing the photographic emulsion layer may be subjected to a bleach/fix process after the bleaching process, or after a fixing process. As a bleaching agent in the bleaching solution or the bleach/fix solution of the present invention, generally use can be made of aminopolycarboxylic acid iron complex salts. As additives used for the bleaching solution or bleach/fix solution of the present invention, use can be made of various compounds described in Japanese Patent Application No. 32462/1986, pages 22 to 30.

When the color developing solution is substantially free from benzyl alcohol, the leucolization reaction of a cyan dye in the bleach/fix solution barely occurs, so that the pH of the bleach/fix solution or the amount of an oxidizing agent can be lowered.

The term "substantially free from benzyl alcohol" means the content of benzyl alcohol is 0.5 m1/1 or below.

The replenishing amount of the bleach/fix solution is generally about 330 m t/m^2 or below, and if the color developing solution does not contain benzyl alcohol the replenishing amount can be lowered to 60 m t/m^2 or below.

After the desilvering step (bleach/fix or fix), water washing and/or stabilizing or the like is carried out. As additives used in the washing and stabilizing steps, use can be made of various compounds described in Japanese Patent Application No. 32462/1986, pages 30 to 36.

It is preferable that the amount of the replenishing solution of each process be smaller. It is preferable that the amount of the replenishing solution is 0.1 to 50 times, more preferably 3 to 30 times, as much as

the carried-over amount from the preceding bath per unit area of the photographic material.

According to the invention, an excellent effect can be exhibited wherein after a silver halide color photographic material has been color-developed, bleached, and fixed, the white background of the color photograph can be prevented from discoloring even during long-term storage or display.

Further, according to the invention, an excellent effect can be exhibited wherein deterioration of a dye image due to the remaining color developing agent taken into the photographic material after the color development, bleaching, and fixing processes, or due to its oxidized product, can be prevented. Still further, according to the invention, an excellent effect can be exhibited wherein even if the color photographic material is processed with a processing solution in a running state, a processing solution that will be washed with less water or will not be washed with water, a processing solution that is substantially free from benzyl alcohol, such as a color developing solution, and whose components will be brought into the photographic material in a greater amount, or other processing solutions that will for example, impose a burden on the color development, image deterioration due to the remaining aromatic amine developing agent or its oxidized product and the occurrence of stain or side effects therefrom can be prevented.

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Examples

Examples of the invention are given below, but the invention is not limited to the examples.

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Example 1

A color photographic material (A-1) was prepared by coatings the first layer (lowermost layer) to the seventh layer (uppermost layer) of the compositions shown in Table 1 on a both-sides polyethylene-laminated paper base.

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Table 1 Integration delatin Actyl-Woodfixed Copolymer of Polyvinyl alcohol (modification degree 17%) 1.33 Colour (a) Solvent (b) Solvent (c)	50	4 5	40 .	35	30		25	20	15	10	5
mage bye Stabilizer (A-30) Silver Chlorobromide Emulsion (silver bromide 75 molf) Silver (A-30) Silver (A-30											•
Galatin Gal	Layer				Main compo	nent					
Gelatin Solvent (4) Solvent (5) Solvent (6) Solvent (7) Solvent (7) Solvent (8) Solvent (9) Solvent (9) Solvent (10) Solve	Seventh Layer (protective layer)	Gelat. Acryl-	odified		f Polyviny	l alcohol	(modifice	ation degr		1.33	8/m²
Silver Chlorobromide Emplsion (silver bromide 70 molf) Silver (b) Gelatin Gelatin Gelatin UV Absorbent (d) Color Mix Inhibitor (A-30) Solvent (a) Solvent (b) Solvent (c) Solvent (c) Solvent (d) Color Mix Inhibitor (A-18) Solvent (d) Gelatin Mageing Dup Stabilizer (A-18) Gelatin Olor Mix Inhibitor (b) Solvent (c) Solvent (d) Gelatin	Sixth Layer (UV ray absorbing layer)	Gelat: UV Abs Solver		. (0.54 5.10 × 10-* 0.08	1
Gelatin UV Absorbent (d) Color Mix Inhibitor (A-30) Silver Chiorobromide Emulsion (silver bromide 75 molf) Silver: Silver Chiorobromide Emulsion (silver bromide 80 molf) Silver: Gelatin Gelatin Gelatin Gelatin Gelatin Gelatin Gelatin Gelatin Silver: Gelatin	Fifth Layer (red-sensitive emulsion layer)	Silver Gelati Cyan (Image Solver	r Chlorobromid in Coupler Dye Stabilize	le Emils: r (f)	ion (silve	r bromide	70 mol#)	-	Silve	1	Ĕ
Silver Chlorobromide Emulsion (silver bromide 75 molf) Silver: 0.15 Hagenta Coupler Image Dye Stabilizer (A-18) Solvent (c) Gelatin Gelatin Gelatin Color Mix Inhibitor (b) Silver Chlorobromide Emulsion (silver bromide 80 molf) Silver (A-43) Solvent (a) Polyethylene Laminate Paper (a white pigment (TiO ₂) and a bluish dye (ultramarine) were included in the first lavar side)	Fourth Layer (UV absorbing layer)	Gelatí UV Abs Color Solven	in Sorbent (d) Mix Inhibitor it (a)	(A-30)						1.70 x 10-4 1.60 x 10-4 0.24	g/m² mol/m² g/m²
Gelatin Color Mix Inhibitor (b) Silver Chlorobromide Emulsion (silver bromide 80 molf) Silver: 0.35 g/m Gelatin Yellow Coupler Image Dye Stabilizer (A-43) Solvent (a) Polyethylene Laminate Paper [a white pigment (TiO2) and a bluish dye (ultramarine) were included in the first layer adde)	Third Layer (green-sensitive emulsion layer)	Silver Gelati Magent Image Solven		e Emulsi	on (silver	bromide.	75 mol#)		S11ve	. 0.15 1.56 3.38 x 10 0.19	8/m² mo1/m² .8/m²
Silver Chlorobromide Emulsion (silver bromide 80 molf) Silver: 0.35 g/m Gelatin Yellow Coupler 1.35 g/m Yellow Coupler 1.35 mol/mise Dye Stabilizer (A-43) 8/mise Dye Dye Dye Dye Dye Dye Dye Dye Dye Dy	Second Layer (color mix preventing layer)	Gelati Color	n Mix Inhibitor			-				0.90 2.33 x 10 -1	g/m² mol/m²
(a white pigment (TiO2) and in the first laver side:	First Layer (blue-sensitive emulsion layer)	Silver Gelati Yellow Image Solven	Chlorobromide n Coupler Dye Stabilizer t (a)	e Emulsi (A-43)	on (silver	bromide	80 mol#)		Silver	"	8/m² "mol/m² 8/m²
		Polyet! (ultran	hylene Laminat marine) were i	e Paper		pigment (1	rio, and		İye		

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As spectral sensitizers for the respective emulsion layers, the following compounds were used.

5 Blue-sensitive Emulsion Layer;

Ce
$$CH_2$$
 CH_2 CH_3 CH_4 CH_2 CH_2 CH_3 CH_4 CH_4 CH_5 CH_5

(2 x 10 4 mol per mol of silver halide)

Green-sensitive Emulsion Layer;

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$$Cl \xrightarrow{O} CH = C - CH = O$$

$$Cl \xrightarrow{O} CH = C - CH = O$$

$$CH_{2})_{3}SO_{3}\Theta$$

$$(CH_{2})_{2}SO_{3}Na$$

(2.5 x 10 4 mol per mol of silver halide)

Red-sensitive Emulsion Layer:

 $CH_3 CH_3$ CH_3 H

(2.5 x 10 4 mol per mol of silver halide)

(a) Solvent

(iso C₉H₁₉O) 3 P=0

(b) Color mix inhibitor

(c) Solvent

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(C8 H1 7 O)3 P=O and
$$\begin{pmatrix} CH_3 \\ - \\ - \end{pmatrix} P = O$$

(mixture in weight ratio of 2:1)

(d) UV Absorber

(d) UV Absorber

$$Cl \longrightarrow N \longrightarrow C_4H_9(t) \longrightarrow C_4H_9(sec)$$

$$C_4H_9(t) \longrightarrow C_4H_9(sec)$$

$$C_4H_9(sec) \longrightarrow C_4H_9(sec)$$

$$CL \longrightarrow N \longrightarrow C_4H_9(t)$$

$$CH_2CH_2COOC_8H_{17}$$

(mixture in molar ratio of 1:5:3)

(e) Solvent

$$\begin{pmatrix}
CH_3 \\
X \\
D
\end{pmatrix} P = O$$

(f) Image Dye Stabilizer

5
$$Cl$$
 N N $C_4H_9(t)$ OH $C_4H_9(t)$ OH $C_4H_9(t)$ and OH $C_4H_9(sec)$ OH $C_4H_9(sec)$ OH $C_4H_9(sec)$

(mixture in molar ratio of 1:3:3)

A - 30

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A - 4 3

(t)C₄H₉

$$(HO \longrightarrow CH_2 + CC + CO_2 \longrightarrow CH_3$$

$$(t)C_4H_9 \longrightarrow CH_2 \longrightarrow CCH_3$$

$$(CH_3 \longrightarrow CCH_3)$$

$$(CH_3 \longrightarrow CCH_3$$

The following dyes were used to protect the respective emulsion layers from irradiation;

Green-sensitive Emulsion Layer;

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Red-sensitive Emulsion Layer;

HOOC NO CH-CH=CH-CH=CH NO COOH

Of these layers, the yellow coupler in the first layer was a mixture of equal mol of the above-mentioned (Y-1) and (Y-7) to obtain a coating amount of 6.91×10^4 mol/m². The cyan coupler in the fifth layer was a mixture of equal mol of the above-mentioned (C-27) and (C-10) to obtain a coating amount of 7.05×10^4 mol/m². The magenta coupler in the third layer was (M-30) to obtain a coating amount of 3.38×10^4 mol/m².

Then Samples (A-2) - (A-12) were prepared by adding the preservative compounds of the present invention in the third layer (green-sensitive layer) of Sample (A-1). In some of these samples the magenta coupler (M-37) was used instead of (M-30). The details of these Samples are shown in Table 2.

Each of the thus prepared Samples was subjected to a exposure through an optical wedge and then processed according to the following processing procedure (I) to obtain a color Image.

Processing Procedure (I)

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A running developing process was carried out in the following steps and conditions using a Fuji Color Roll Processor FMPP 1000 (partiary reconstructed)(processor made by Fuji Photo Film Co.).

10	Step	Time (sec.)	Tempera- rature (°C)	Tank Capacity (1)	Replenisher amount (m2/m²)*
	Color developing	45	35	88	150
15	Bleach-fixing	45	35	35	50
	Rinsing ①	20	35	17	-
20	Rinsing ②	20	35	17	-
	Rinsing ③	20	35	17	250

^{*} per m² of the photographic material

The rinsing steps were carried out in a three-tank counter-current mode, in which the replenisher is fed to tank of rinsing 3, the overflow rinsing solution from tank of rinsing 3 is fed to the bottom of rinsing tank of rinsing 2, the overflow rinsing solution from tank of rinsing 3 is fed to the bottom of rinsing tank of rinsing 1, and the overflow rinsing solution from tank of rinsing 1 is drained off. The carried-over amount of solution from each tank was 25 mL/m^2 of paper.

The composition of each tank solution and replenisher were as follows:

Color Developing Solution

35	,	Tank Solution	Replenisher
	Water	800 ml	800 mg
40	Diethylenetriaminepenta-		
	acetate	3.0 g	3.0 g
	Benzyl alcohol	15 ml	17 mg
45	Diethyleneglycohol	10 mg	10 mg
	Sodium sulfite	2.0 g	2.5 g
50	Potassium bromide	0.5 g	
	Sodium carbonate	30 g	30 g
	N-Etyl-N-(β-methanesulfonamido-		
55	ethyl)-3-methyl-4-aminoaniline		•
	sulfonate	5.0 g	7.0 g

	. *						
••	Hydroxylamine sulfonate	4.0 g	4.5 g				
10	Brightening agent	1.0 g	1.5 g				
	Water to make	1000 m£	1000 mL				
15	pН	10.10	10.50				
	Bleach-fixing Solution						
20		Tank Solution	Replenisher				
	Water	400 m2	400 ml				
25	Ammonium thiosulfite						
20	(70% solution)	150 mL	300 ml				
	Sodium sulfite	12 g	25 g				
30	Ammonium iron (III) ethylene-	-					
	diaminetetraacetate	55 g	110 g				
25	Disodium ethylenediaminetetra	acetate 5 g	10 g				
35	Water to make	1000 mL	1000 ml				
	рН	6.70	6.50				
40	Rinsing Solution						
	Ethylenediamine-N,N,N',N'-tet	ra-					
45	methylene phosphonate	0.3	g				
~	Benzotriazole	1.0 g					
	Water to make	1000 n	n &				
50	pH (by sodium hydroxide)	7.	5				

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Processing Procedure (II)

5	Step	Time	Tank Capacity (1)	Replenisher Amount (mg/m²)*
•	Color developing	45 sec.	88	150
10	Bleach-fixing	2 min. 0 sec.	35	_~ 350
	Rinsing $\textcircled{1}$	1 min. 0 sec.	17	_
15	Rinsing ②	1 min. 0 sec.	17	-
	Rinsing ③	l min. O sec.	17	1,300

^{*}per m^2 of the photographic material

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Processing solutions and replenishers having the same compositions as the processing procedure (I), respectively, were used.

At the point of one hour after development processing according to the above-mentioned procedure, a magenta reflective density was measured at a non-image area of each processed sample of photographic material. The same measurements were carried out again on the processed samples after being kept for 7 days at 80°C and 10 - 15% RH, and on the processed samples after being kept for 8 days at 80°C and 70% RH. The results are shown in Table 2 in values of increments of stain after one hour.

5	-	Remarks		Comparative) = 1	s	=	=	: `	=	#	This	=	=	Comparative	E .	Þ	E	I	This	, , , , , , , , , , , , , , , , , , ,	Ľ	=	
10		of Magenta Stain	8 days at 80°C, 70% RH	0.02	60.0	0.08	0.03	20.0	0.11	0.15	0.08	0.02	0.01	0.01	0.19	0.08	80.0	0.03	0.11	0.02	10.0	0.02	0.01	
15		Increment of Ma	/B at	90.0	0.04	0.02	10.0	0.02	0.04	0.05	0.03	0.01	0.01	0.01	0.05	0.04	0.03	0.02	0.03	0.01	0.01	0.01	0.01	
20			7 days 80 c	0	·	ó	o _	σ .	ŏ	ó	0		ó	•	0	0	0	0	٥.	0	0	Ó	0	
25	Table 2	Processing	Procedure	(1)	(11)	(1)	(11)	(I)	(I)	(I)	(I) ·	(1)	(I)	(1)	(1)	(11)	(I)	(I)	Œ	(I)	(I)	(1)	(I)	
30		Amount of	Additive (molf to coupler)	-		20 ,	50	. 02	20	20	20	10 + 10	10 + 10	10 + 10		,	20	20	20	10 + 10	10 + 10	10 + 10	10 + 10	
35		, as	id)				_				8)	(1-1)	(2-1	1-18)				-	1)	1-21)	(111-21)	1-21)	(III-18)	
40		Additiv	(Exemplified Compound)			(h-I)		(1-27)	(II-5)	(2-III)	(111-18)	(L-III) + (h-I)	(1-27)+ (111-7)	(II-5)+ (III-18)			(I-22)	(I-30)	(111-21)	(I-22)+ (III-21)	(r-27)+ (II:	(I-30)+ (III-2I)	(I-22)+ (II	
45		Magents	Coupler	M - 30	=	E	z	3	=	5	=	3	=	z	М - 37	2	ε		E	±	=	E	£	
50	**	Samole		A - 1	A - 3	A - 2	A - 2	A - 3	- ¥	A - 5	A - 6	A - 7	80 1 W	6 • V	A -10	A -10	A -11	A -12	A -13	A -14	A -15	A -16	A -17	

As is apparent from the results in Table 2, the increments of magenta stain are relatively small on the samples processed according to the procedure (II) in which the bleaching time and rinsing times were longer and sufficient amounts of replenisher were used, but magenta stain was greatly increased on samples processed by the procedure (I) in which the processing times were shorter and the replenisher amounts were smaller.

From the results described above, it can been seen that satisfactory prevention of magenta stain is

possible by the combined use of the preservability improving compounds (A) and (B) of the present invention, although the prevention is not sufficient on samples in which compounds (A) and (B) were used separately.

Compounds to be used in Examples 2 - 8 are as follows:

Sensitizing dye

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ExS-1

$$C\ell \xrightarrow{S} CH \xrightarrow{S} C\ell$$

$$(CH_2)_4 SO_3^{\Theta} (CH_2)_4$$

$$SO_3 HN(C_2H_5)_3$$

ExS-2

$$\begin{array}{c|c}
C & C_2H_5 \\
C & C_1H_5 \\
C & C_1H_5 \\
C & C_2H_5 \\
C & C_1H_5 \\
C & C_1H_2 \\
C & C_1H$$

ExS-3

$$\begin{array}{c} O \\ \bigoplus \\ N \\ (CH_2)_4 SO_3 \\ \Theta \\ (CH_2)_4 \\ SO_3 \\ HN(C_2H_5)_3 \end{array}$$

ExS-4

$$C\ell \xrightarrow{S} CH \xrightarrow{S} CH \xrightarrow{N} CH \xrightarrow{N} CH_{2})_{3}$$

$$(CH_{2})_{4}SO_{3}^{\Theta} \xrightarrow{(CH_{2})_{3}} (CH_{2})_{3}$$

$$SO_{3}HN(C_{2}H_{5})_{3}$$

ExS-5

$$\begin{array}{c|c}
C_2H_5 \\
O \\
CH=C-CH = \\
N \\
(CH_2)_2SO_3\Theta \\
(CH_2)_2
\end{array}$$

ExS-6

$$\begin{array}{c|c}
O & S \\
N & N \\
CH_2 & CH_2
\end{array}$$

$$\begin{array}{c|c}
CH_2 & S \\
CH_2 & S \\
N(C_2H_5)_3
\end{array}$$

ExS-7

$$\begin{array}{c} C_2H_5 \\ C_2H_5 \\ C_1 \\ C_2 \\ C_3 \\ C_4 \\ C_5 \\ C_4 \\ C_5 \\ C_6 \\ C_7 \\ C_8 \\$$

ExS-8

$$\begin{array}{c|c}
CH_3 & CH_3 \\
CH & CH \\
CH & CH
\end{array}$$

$$\begin{array}{c|c}
CH_3 & CH_3 \\
CH & CH
\end{array}$$

ExS-9

$$\begin{array}{c|c}
CH_3 & CH_3 & CH_3 \\
CH_3 & CH_2 & CH_3
\end{array}$$

$$\begin{array}{c}
CH_3 & CH_3 \\
CH_2 & CH_3
\end{array}$$

$$\begin{array}{c}
CH_3 & CH_3 \\
CH_2 & CH_3
\end{array}$$

$$\begin{array}{c}
CH_3 & CH_3 \\
CH_2 & CH_3
\end{array}$$

$$\begin{array}{c}
CH_3 & CH_3 \\
CH_2 & CH_3
\end{array}$$

$$\begin{array}{c}
CH_3 & CH_3 \\
CH_2 & CH_3
\end{array}$$

ExS-10

$$\begin{array}{c|c}
S & C_2H_5 \\
\downarrow & \downarrow \\
N & CH=C-CH = \\
N & \downarrow \\
N & CH_2
\end{array}$$

$$\begin{array}{c|c}
C_2H_5 & S \\
\downarrow & \downarrow \\
CH_2
\end{array}$$

$$\begin{array}{c|c}
C_1H_5 & S \\
\downarrow & \downarrow \\
CH_2
\end{array}$$

$$\begin{array}{c|c}
C_1H_2 & S_3HN(C_2H_5)_3
\end{array}$$

ExS-11

$$C\ell \xrightarrow{S} CH = C - CH = C$$

$$C\ell \xrightarrow{S} CH = C - CH = C$$

$$C\ell \xrightarrow{N} C\ell$$

ExS-12

10

15

20

ExY-1

 $\begin{array}{c|c} CH_3 & C\ell \\ CH_3 - C - COCHCONH - C \\ CH_3 & NHCOCHO - C_5H_{11}(t) \\ O = C & C_2H_5 \\ CH_3 - O & CH_3 \end{array}$

ExY-2

25

$$\begin{array}{c|c} CH_3 & C\ell \\ CH_3 - C - COCHCONH - \\ CH_3 & NHCOCHO - C_5H_{11}(t) \\ CH_3 & C_2H_5 \\ C_2H_5 & C_2H_5 \end{array}$$

35

30

⁴⁰ ExY-3

CH₃ Cl CH₃ C C COCHCONH NHCO-(CH₂)₃ O C₅H₁₁(t) CH₃ NHCO-(CH₂)₃ O C₅H₁₁(t) NHCO-(CH₂)₃ O C₅H₁₁(t)

ExM-1

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· 45

 $CH_3 \qquad C\ell$ $N \qquad NH \qquad OC_8H_{17}$ $CHCH_2NHSO_2 \qquad OC_8H_{17}$ $CH_3 \qquad NHSO_2 \qquad C_8H_{17}(t)$

ExM-2

CH₃ Cl N NH OCH₂CH₂OCH₂CH₃ N= $\begin{pmatrix} CHCH_2NHSO_2 & OC_8H_{17} \\ CH_3 & NHSO_2 & C_8H_{17} \end{pmatrix}$

ExM-3

 OC_4H_9 OCH_3 OCH_3 $NHSO_2$ OC_8H_{17} $NHSO_2$ OC_8H_{17} $NHSO_2$

55

10

15

Ex M-4

20

25

ExM-5

30

(n) C_6H_{13} CHCH₂ SO₂CH₂CH₂

35

ExM-6

40

 $\begin{array}{c|c}
 & Ct \\
 & N

Ć₈H₁₇(t)

 $VHSO_2 - C_8H_{17}(t)$ $C_8H_{17}(t)$

50

$$ExC-1$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{11}$$

20 ExC-2

$$C_{2}H_{5} \xrightarrow{OH} NHCOCHO \xrightarrow{C_{4}H_{9}} (t)C_{5}H_{11}$$

ExC-3

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

ExC-4

$$Cl \longrightarrow NHCOCHO \longrightarrow (t)C_5H_{11}$$

$$CH_3 \longrightarrow (t)C_5H_{11}$$

ExC-5

$$(t)C_5H_{11} \longrightarrow \begin{matrix} C_6H_{13} \\ OCHCONH \end{matrix} \longrightarrow \begin{matrix} C_\ell \\ C_\ell \end{matrix}$$

ExC-6

ExC-7

$$C_{2}H_{5}$$

$$C_{2}H_{5}$$

$$C_{3}H_{11}(t)$$

$$C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t)$$

ExC-8

$$(t)C_5H_{11} \longrightarrow O - CHCONH \longrightarrow F F$$

$$C_5H_{11}(t) \longrightarrow CL$$

$$(t)C_5H_{11}(t) \longrightarrow F$$

$$C_7H_{11}(t) \longrightarrow F$$

Cpd-1

$$\begin{array}{c|c}
Cl & OH & C_4H_9(t) \\
\hline
 & N & C_4H_9(t)
\end{array}$$

$$\begin{array}{c}
C_4H_9(t)
\end{array}$$

Cpd-2

Cpd-3

Cpd-4

Cpd-5

$$\begin{array}{c}
C_4H_9(t) & O \\
HO \longrightarrow CO \longrightarrow C_4H_9(t)
\end{array}$$

Cpd-6

$$(t)C_8H_{17} \xrightarrow{OH} C_8H_{17}(t)$$

Cpd-7

(sec)
$$C_8H_{17}$$
 OH OH OH

$$Cpd-8$$

$$-(CH_2-CH_{\frac{1}{n}})$$
 (n=100~1000)
CONHC₄H₉(t)

45 Cpd-10

Poly(ethyl acrylate)latex

Cpd-11

$$Cpd-12$$

$$\begin{pmatrix}
(t)C_4H_9 \\
HO - CH_2 \\
(t)C_4H_9
\end{pmatrix}$$

$$CH_3 CH_3 O \\
N - CCH = CH_2$$

$$CH_3 CH_3$$

$$CH_3 CH_3$$

$$Cpd-13$$

Cpd-14

5 Cpd-15 -CH-CH=CH-CH=CH 10 15 SO₃K Cpd-16H₅C₂OOC-7 CH-CH=CH-CH=CH T COOC₂H₅ 20 HO ÇH₂ 25 SO₃K SO₃K Cpd-17 30 HO OH || | CNCH₂CH₂OH HOCH2CH2NC 35 SO₃Na SO₃Na 40 Cpd-18HO HOCH₂CH₂NC CNCH2CH2OH 45 N I CH₂ ĊH₂ 50

114

SO₃Na

55

SO₃Na

Cpd - 19

5 CH₃ N N N OH

Cpd - 20

10

30

Cpd-21

Cpd-22

NaO-C
$$CH$$
-CH=CH CH -CH=CH CH
N O HO
N O
C-ONa

CH₂
SO₃Na
SO₃Na

Solv-1: Di(2-ethylhexyl)phthalate

Solv-2: Trinonylphosphate

Solv-3: Di(3-methylhexyl)phthalate

Solv-4 : Tricresylphthalate Solv-5 : Dibutylphthalate

Solv-6: Trioctylphosphate

Solv-7: Diethylazelate

Solv-8: Dioctylsebacate

Example 2

A color photographic material (B-1) was prepared by multi-coatings composed of the first to the twelfth layer as hereinbelow defined and coated on a both-sides polyethylene-laminated paper base. A white pigment (TiO2) and a small amount of bluish dye (ultramarine blue) were included in the first layer side of the polyethylene film laminated.

Composition of photosensitive layers

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In the following compositions, each ingredient is indicated in g/m2 of a coating amount, but the coating amount of the silver halide is shown in g/m2 in terms of silver.

15

First layer: Geratin layer Gelatin

1.30

Second layer: Antihalation layer Black colloidal silver

Gelatin

0.70

Third layer: Red-sensitive emulsion (low sensitivity) layer Silver chloroiodobromide emulsion spectralsensitized by red-sensitizing dye (ExS-7, -11 and -12) (silver chloride : 1 mol%, silver iodide : 4 mol%, average grain size : 0.3 μm, grain size distribution : 10 %, cubic, core-shell type of iodide core) Silver iodobromide emulsion spectral-sensitized by red-sensitizing dye (ExS-7, -11 and -12) (silver iodide : 5 mol%, average grain size : 0.45 μm , grain size distribution : 20%, plate (aspect ratio:5)) Gelatin 1.00

Cyan coupler (ExC-2) 0.14

Cyan coupler (ExC-5) 0.07

Discoloration inhibitor (equal amount mixture of Cpd-1, -3, -5 and -11) 0.12

Dispersion medium for coupler (Cpd-9) 0.03

Solvent for coupler (Solv-1, -2 and -3) 0.06

35

Fourth layer : Red-sensitive emulsion (highly sensitive) layer Silver iodobromide emulsion spectralsensitized by red-sensitizing dye (ExS-7, -11 and -12) (silver iodide: 6 mol%, average grain size : 0.75 μm, grain size distribution: 25 %, plate (aspect ratio: 8, core-shell type of iodide core) Gelatin 1.00

Cyan coupler (ExC-2) 0.20

> Cyan coupler (ExC-5) 0.10

Discoloration inhibitor (equal amount mixture of Cpd-1, -3, -5 and -11)

Dispersion medium for coupler (Cpd-9) 0.03

Solvent for coupler (Solv-1, -2 and -3) 0.10

45

Fifth layer: Intermediate layer Magenta colloidal silver 0.02

Gelatin 1.00

Color mix inhibitor (Cpd-6 and -13)

Solvent for color mix inhibitor (Solv-4 and -5) 0.16

Polymer latex (Cpd-10) 0.10

Sixth layer: Green-sensitive emulsion (low sensitivity) layer Silver chloroiodobromide emulsion spectralsensitized by green-sensitizing dye (ExS-7)(silver chloride: 1 mol%, silver iodide: 2.5 mol%, average grain size: 0.28 μm, grain distribution: 12%, cubic, core-shell type of iodide core) Silver iodobromide emulsion spectral-sensitized by green-sensitizing dye (ExS-7)(silver iodide : 2.8 mol%, average grain size: 0.45 µm, grain size distribution: 12%, plate (aspect ratio:5))

Gelatin 0.80

Magenta coupler (ExM-1) 0.10

Discoloration inhibitor (Cpd-11) 0.10

Stain inhibitor (Cpd-8) 0.001

Dispersion medium for coupler (Cpd-9) 0.05

Solvent for coupler (Solvent-4 and -6) 0.15

Seventh layer: Green-sensitive emulsion (highly sensitive) layer Silver iodobromide emulsion spectralsensitized by green-sensitizing dye (ExS-7)(silver iodide: 3.5 mol%, average grain size: 0.9 µm, grain size distribution: 23%, plate (aspect ratio: 9, uniform iodide type)) 0.10 Gelatin 0.80

Magenta coupler (ExM-1) 0.10 Stain inhibitor (Cpd-8) 0.001

Dispersion medium for coupler (Cpd-9)
 Solvent for coupler (Solv-4 and -6)
 0.15

Eighth layer: Yellow filter layer Yellow colloidal silver 0.20
Gelatin 1.00
Color mix inhibitor (Cpd-6) 0.06
Solvent for color mix inhibitor (Solv-4 and -5) 0.15

0.10

25

Ninth layer: Blue-sensitive emulsion (low sensitivity)layer Silver chloroiodobromide emulsion spectral-sensitized by blue-sensitizing dye (ExS-5 and -8) (silver chloride: 2 mol%, silveriodobromide: 2.5 mol%, average grain size: $0.35~\mu m$, grain size distribution: 8%, cubic, core-shell type of iodide core) 0.07 Silver iodobromide emulsion spectral-sensitized by blue-sensitizing dye (ExS-5 and -6)(silver iodobromide: $0.56~\mu m$) (silver iodobromide) (silver iodobromide)

30 2.5 mol%, average grain size : 0.45 μm, grain size distribution : 16%, plate (aspect ratio : 6) 0.10 Gelatin 0.50

Yellow coupler (ExY-2) 0.20 Stain inhibitor (Cpd-8) 0.001

Polymer latex (Cpd-10)

Discoloration inhibitor (Cpd-12) 0.10

35 Dispersion medium for coupler (Cpd-9) 0.05 Solvent for coupler (Solv-2) 0.05

Tenth layer: Blue-sensitive emulsion (highly sensitive) layer Silver iodobromide emulsion spectral-sensitized by blue-sensitizing dye (ExS-5 and -6)(silver iodide: 2.5 mol%, average grain size: 1.2 µm, grain size distribution: 21%, plate (aspect ratio:14)) 0.25

Gelatin 1.00 Yellow coupler (ExY-2)

Yellow coupler (ExY-2) 0.40 Stain inhibitor (Cpd-8) 0.002

Discoloration inhibitor (Cpd-12) 0.10
 Dispersion medium for coupler (Cpd-9)0.05
 Solvent for coupler (Solv-2) 0.10

Eleventh layer: UV absorbing layer Gelatin
UV absorbent (Cpd-1, -3 and -4)
Color mix inhibitor (Cpd-6 and -7)
Solvent for UV absorbent (Solv-1 and -2)
Irradiation preventing dye (Cpd-13 and -14)
Irradiation preventing dye (Cpd-15 and -16)
UV absorbent (Solv-1 and -2)
Irradiation preventing dye (Cpd-15 and -16)
UV absorbent (Cpd-13 and -14)
UV absorbent (Cpd-15 and -16)

Twelfth layer: Protective layer Fine grain size silver chlorobromide emulsion (silver chloride: 97 mol%,

average grain size : 0.2 µm) 0.07 Modified polyvinyl alcohol 0.02

Gelatin 1.50

s Sodium 1-oxy-3,5-dichloro-s-triazine 0.17

In addition, Alkanol SC (tradename, made by Dupont) and sodium alkylbenzenesulfonate were used as auxiliary agents for emulsification and dispersion, and succinate ester and Magefac F-120 (tradename, made by Dainippon Ink) were added as coating aids to each layer. Further, Cpd-19, -20 and -21 were used as stabilizers for the layers containing silver halide or colloidal silver.

Samples (B-2) and (B-3) were prepared by repeating the preparation procedure of Sample (B-1), except that magenta coupler (ExM-1) was changed to equal mol of (Exm-2) and (Exm-3) respectively. Then, Samples (B-4) to (B-11) were prepared by adding the preservability improving compounds (A) and/or (B) of the present invention to the sixth layer and the seventh layer of Samples (B-1) to (B-3). The details of the addition of the preservability improving compounds are shown in the following Table 3.

Each of the thus prepared samples was subjected to a exposure through an optical wedge and then to a color development process according to the processing procedure (III) described below.

20 Processing Procedure (III)

	Step	Temperature	Time
25	First developing (Black		
	and white developing)	38 °C	l min. 15 sec.
30	Water-washing	38 °C	1 min. 30 sec.
	Reversal exposure	over 100 Lux	over 1 min.
	Color developing	38°C	2 min. 15 sec.
35	Water washing	38°C	45 sec.
	Bleach-fixing	38°C	2 min.
40	Water washing	38°C	2 min. 15 sec.

Composition of processing solution

45	First Developing Solution
	Pentasodium nitrilo-N,N,N-trimethylenephosphonate 0.6 g
	Pentasodium diethylenetriaminepentaacetate 4.0 g
	Potassium sulfite 30.0 g
	Potassium thiocyanate 1.2 g
50	Potassium carbonate 35.0 g
	Potassium hydroquinonemonosulfonate 25.0 g
	Diethyleneglycol 15.0 m.t.
	1-Phenyl-4-hydroxymethyl-4-methyl-3-pyrazolidone 2.0 g
	Potassium bromide 0.5 g
55	Potassium iodide 5.0 mg
	Water to make 1000 m.t
	(pH 9.70)

0 277 589

Color Developing SolutionBenzyl alcohol 15.0 m L Diethylene glycol 12.0 m t 3,6-Dithia-1,8-octanediol 0.2 gPentasodium nitrilo-N,N,N-trimethylenephosphonate 0.5 gPentasodium diethylenetriaminepentaacetate 2.0 g Sodium sulfite Sodium carbonate 25.0 g Hydroxylamine sulfonate 3.0 g N-Ethyl-N-(β -methanesulfonamidoethyl)-3-methyl-4-aminoaniline sulfonate 5.0 g Potassium bromide 0.5 g Potassium iodide 1.0 mg Water to make 1000 mt (pH 10.40)

15

Bleach-fixing Solution2-Mercapto-1,3,4-triazole 1.0 g
Disodium ethylenediaminetetraacetate dihydrate 5.0 g
Ammonium iron(III) ethylenediaminetetraacetate monohydrate 80.0 g
Sodium sulfite 15.0 g
Sodium thiosulfate (700 g/t solution) 160.0 mt
Glacial acetic acid 5.0 mt
Water to make 1000 mt
(pH 6.50)

At the point of one hour after development processing according to the above-mentioned procedure, a magenta reflective density (stain) was measured at a non-image area of each processed sample of photographic material. The same stain measurements were carried out again on the processed samples after being kept for 3 days at 80°C and 70% RH, and on the processed samples after being kept for 100 days at room temperature. The results are shown in Table 3 in values of increments of stain after one hour.

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	-	Table 3			
- 1	44444	Amount of Additive	Increment of Magenta Stain	agenta Stain	Remarks
	(Exemplified Compound)	(mol% to coupler)	3 days at 80 C, 70%RH	100 days at R.T.	
1	-		0.08	0.07	Comparative Example
			0.07	0.05	
			60.0	0.08	t
	(1-26)	20	0.03	0.02	=
	(I-26) + (III-19)	10 + 10	0.01	00.00	This
	(1-5)	50	0.03	0.02	Comparative
	(L-III)	20	0.05	0.03	exampre "
	(1-2) + (111-1)	10 + 10	0.01	0.01	This
	(II-3)	20	0.07	0.05	Comparative
	(III-24)	50	0.04	0.03	exampte
	(11-3) + (111-24)	10 + 10	0.01	0.01	This
			,		

As is apparent from the results in Table 3, it can be understood that the stain-preventive effect according to the present invention is quite remarkable, and it can also be seen that this effect does not decline even if the structure of photographic material and the development processing solutions are varied.

Example 3

A multi-layer color photographic paper (C-1) was prepared which has such layers as hereinbelow described on a paper laminated on both sides with polyethylene. Coating solutions were prepared as follows:

Preparation of the first layer coating solution

To a mixture of 10.2 g of yellow coupler (ExY-1), 9.1 g of yellow coupler (ExY-2) and 4.4 g of a image dye stabilizer (Cpd-12), 27.2 mt of ethyl acetate and 7.7 mt (8.0 g) of high boiling solvent (Solv-5) were added, and they were dissolved. The resulting solution was emulsified and dispersed in 185 mt of 10% aqueous gelatin solution containing 8 m t of a 10% solution of sodium dodecylbenzenesulfonate. Each of the under-mentioned emulsions EM1 and EM2 was mixed with the above-obtained emulsified and dispersed solution and dissolved, and the concentration of gelatin in the mixture was adjusted so as to obtain the composition shown below, thereby preparing the first coating solution. The second to the seventh layer coating solutions were prepared in the same maner as the first coating solution. As a gelatin hardner for the respective layers, the sodium salt of 1-oxy-3,5-dichloro-s-triazine was used.

20

Compositions of layers

The composition of each layer is shown below. Each ingredient is indicated in g/m² of a coating amount, but the coating amount of silver halide is shown in g/m2 in terms of silver.

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Supporting base

Polyethylene laminated paper (a white pigment, TiO2 and a bluish dye, ultramarine, were included in the first layer side of the polyethylene film laminated).

First layer: Blue-sensitive layer Monodisperse silver chlorobromide emulsion (EM1) spectral-sensitized by sensitizing dye (ExS-1)

Monodisperse silver chlorobromide emulsion (EM2) spectral-sensitized by sensitizing dye (ExS-1) 0.13

Gelatin 1.86

Yellow coupler (ExY-1) 0.44

Yellow coupler (ExY-2) 0.39

Image dye stabilizer (Cpd-12) 0.19

Solvent (Solv-5) 0.35

Second layer: Color mix preventing layer Gelatin 0.99

Color mix inhibitor (Cpd-7) 0.08

Third layer: Green-sensitive emulsion layer Monodisperse silver chlorobromide emulsion (EM3) spectralsensitized by sensitizing dye (ExS-2,-3) 0.05

Monodisperse silver chlorobromide emulsion (EM4) spectral-sensitized by sensitizing dye (ExS-2,-3)

0.11

Gelatin

Magenta coupler (ExM-1)

Image dye stabilizer (Cpd-11) 0.20

Solvent (Solv-4) 0.12

Solvent (Solv-6) 0.25 Fourth layer: UV absorbing layer Gelatin 1.60

UV absorbent (Cpd-1/Cpd-2/Cpd-3 = 3/2/6 in wt. ratio 0.70

Color mix inhibitor (Cpd-6) 0.05

Solvent (Solv-2) 0.27

5

Fifth layer: Red-sensitive emulsion layer Monodisperse silver chlorobromide emulsion (EM5) spectral-sensitized by sensitizing dye (ExS-8,-12) 0.07

Monodisperse silver chlorobromide emulsion (EM6) spectral-sensitized by sensitizing dye (ExS-8,-12)

10 0.16

Gelatin 0.92

Cyan coupler (ExC-6) 0.32

Image dye stabilizer (Cpd-2/Cpd-3/Cpd-4 = 3/4/2 in wt. ratio) 0.17

Polymer for dispersion (Cpd-9) 0.28

15 Solvent (Solv-4) 0.20

Sixth layer: UV absorbing layer Gelatin 0.54

UV absorbent (Cpd-1/Cpd-3/Cpd-4 = 1/5/3 in wt. ratio) 0.21

o Solvent (Solv-4) 0.08

Seventh layer: Protective layer Gelatin 1.33

Acryl-modified copolymer of polyvinyl alcohol (modification degree: 17%) 0.17

25 Liquid paraffin 0.03

For preventing irradiation, the dyes (Cpd-1 and -2) were used.

Additionally, the same auxiliary agents for emulsification and dispersion, and coating aids as in Example 2 were used. As the stabilizer of silver halide, (Cpd-19) and (Cpd-21) were used. The silver halide emulsion used in this Example were as follows:

	Emulsion	Shape	Grain size (μṃ)	Br Content (mol %)	Fluctuation coefficient.
	EM1	Cubic	1.0	80	0.08
	EM2	Cubic	0.75	80	0.07
1	EM3	Cubic	0.5	83	0.09
	EM4	Cubic	0.4	83	0.10
	EM5	Cubic	0.5	73	0.09
	ЕМ 6	Cubic	0.4	73	0.10

Next, Samples (C-2) to (C-16) were prepared by exchanging the magenta coupler in the third layer (green-sensitive layer) of Sample (C-1) and/or by adding the preservability improving compound of the present invention. The details of the Samples are shown in Table 4.

Each of the thus prepared samles were subjected to a exposure through an optical wedge and then to a processing procedure (IV) decribed below using a Fuji Color Paper Processor FPRR 115 (processor made by Fuji Photo Film Co.) to obtain a color image.

Processing Procedure(IV)

Step	Temp.	Time	Replenisher Amount (ml/m²)*	Tank Capacity (l)
Color developing	37	3 min.30 sec.	200	60
Bleach-fixing	33	1 min.30 sec.	55	40
Water washing ①	24-34	1 min.	-	20
Water washing ②	24-34	l min.	-	20
Water washing ③	24-34	l min.	10	20
Drying	70-80	l min.		

^{*} per m² of the photographic material

The compositions of the processing solution were as follows:

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Color Developing Solution

		Tank Solution	Replenisher
400	Water	800 ml	800 ml
45	Diethylenetriaminepentaacetate	1.0 g	1.0 g
	Nitolirotriacetic acid	2.0 g	2.0 g

50

Water washing steps were carried out in a three-tank cascade mode from tank of washing (3) toward tank of washing (1).

	Benzyl alcohol	15	m L	23	m l
	Diethylene glycol	10	mL	10	m L
, 5	Sodium sulfite	2.0	g	3.0	g
	Potassium bromide	1.2	g	-	
10	Potassium carbonate	30	g	25	g
	N-Ethyl-N-(β-methanesulfonamidoet	nyl)-			
15	3-methyl-4-aminoaniline sulfonate	5.0	g	9.0	g
	Hydroxylaminesulfonate	3.0	g	4.5	g
	Brightening agent (WHITEX4B, trade	ename,	made	by Sum	itomo
20	Chemical)	1.0	g	2.0	g
	Water to make	1000	m L	1000	m L
25	pH (at 25°C)	10.2	20	10.8	80

Bleach-fixing Solution

. 55

30	`	Tank Solution	Replenisher
	Water	400 m &	400 m g
35	Ammonium thiosulfate (70% solution	on)150 ml	300 ml
	Sodium sulfite	13 g	26 g
	Ammonium iron (III) ethylenediam	ine-	
40	tetraacetate	55 g	110 g
•	Disodium ethylenediaminetetraace	tate 5 g	10 g
45	Water to make	1000 ml	1000 ml
	pH (at 25°C)	6.70	6.30

At the point of one hour after development processing by the processing procedure (IV), a magenta reflective density (stain) was measured at a non-image area of each processed sample. The same stain measurements were carried out again on the processed samples after being kept for 14 days at 60°C and 70% RH, and on the processed samples after being kept for 100 days at room temperature. The results are shown in Table 4 in values of increments of stain after one hour.

5		Remarks		Comparative	evampro evampro	=	s	=	:	This	Comparative	ביאפווים ו	This	Comparative	מילוווים ו	This	Comparative	a Tourn	This Invention	
10		Magenta Stain	100 days at R.T.	0.11	0.10	0.03	0.11	0.04	0.03	0.01	0.04	0.03	0.01	0.02	0.02	0.01	0.04	0.02	0.01	
15 20	·	Increment of Ma	14 days at 60°C, 70%RH	0.12	0.10	0.05	0.11	0.05	40.0	0.02	0.04	40.0	0.01	0.03	0.02	10.0	0.05	0.03	0.01	
25	Table 4	Amount of Additive	to coupler)					20	20	10 + 10	20	20	10 + 10	20	20	10 + 10	20		10 + 10	
30		Amount	(mol%	-				-		ਜੋ			Ä			7			7	
35		Additive	(Exemplified Compound)	-				(I-18)	(111-19)	(61-III) + (81-I)	(1-26)	(111-7)	(I-26) + (III-7)	(11-3)	(III-4)	(II-3) + (III-4)	(1-5)	(111-19)	(1-5) + (111-19)	
40										<u>.</u>			<u>.</u>		<u></u>	IJ —		<u> </u>	U —	
45	-	Magenta	Coupler	ExM-1	ExM-2	ExM-3	ExM-4	ExM-1	=	=	ExM-2	=		ExM-3	F	E	ExM-4	E	*.	
50		Sample	•	C - 1	0 - 2	C - 3	# · O	0 - 5	9 - 0	c - 7	8 - O	6 - 0	c -10	c -11	C -12	c -13	C -14	C -15	c -16	

As is apparent from the results in Table 4, it can be understood that the object of the present invention can be attained by the combined use of the preservability improving compounds (A) and (B) of the invention, although magenta stain was not sufficiently prevented by the individual use of compound (A) or 55

Example 4

In a manner similar to Example 1 and Example 3, each of the photographic samples (A-1) to (A-17) of Example 1 and (C-1) to (C-16) of Example 3 was subjected to a exposure through an optical wedge. It was then processed according to the following procedure (V) to obtain a color image.

Processing Procedure(V)

1	ì	ľ.		
•	c	_	•	

	Step	Temperature (°C)	Time
15	Color developing	38	l min. 40 sec.
	Bleach-fixing	30-34	1 min.
	Rinsing (1)	30-34	20 sec.
20	Rinsing ②	30-34	20 sec.
	Rinsing ③	30-34	20 sec.
25	Drying	70-80	50 sec.

Rinsing steps were carried out in a three-tank countercurrent mode from tank of rinsing 3 towards tank of rinsing (1). 30

5.5 g

The composition of the processing solutions were as follows:

Color Developing SolutionWater

800 mt

Diethylenetriaminetentaacetate

1.0 g

1-Hydroxyethylidene-1,1-diphosphonate (60%)

Nitolirotriacetic acid

2.0 g

1,3-Diamino-2-propanol

6.0 g

1,4-Diazabicyclo [2,2,2] octane Potassium bromide

0.5 g

Potassium carbonate 30 g

N-Ethyl-N-(β-methanesulfonamidoethyl)-3-methyl-4-aminoaniline sulfonate

Hydoxylamine sulfonate 4.0 g

Brightening agent (UVITEX, tradename, made by Ciba-Geigy)

Water to make 1000 mL

pH (at 25°C) 10.25

Bleach-fixing SolutionWater 400 mt

Ammonium thiosulfate (70% solution) 200 mt

Sodium sulfite 20 g

Ammonium iron(III) ethylenediaminetetraacetate 60 g

Disodium ethylenediaminetetraacetate

Water to make 1000 mt

> pH (at 25°C) 7.00

Rinsing Solution

Ion-exchanged water (containing under 3 ppm of Ca and Mg, respectively)

Then, as in Example 3, magenta reflective density (stain) measurements were taken at a non-image area on the processed samples after one hour lapsed from the development processing, on the processed samples after being kept for 14 days at 60°C and 70% RH, and on the processed samples after being kept for 100 days at room temperature, respectively. From the results of evaluating the values of increments of magenta stain after one hour, stain increments were not substantially or at all observed on each sample that used preservability improving compounds (A) and (B) of the present invention in combination, although the stain of comparative samples that used compounds (A) and (B) separately did increase.

Example 5

A photographic material (D-1) was prepared in the same manner as described in Sample (C-1) of Example 3 except that silver halide emulsions (EM7-EM12) were used instead of silver halide emulsions (EM1-EM6) respectively.

Emulsion	Shape	Grain size(µm)	Cl Contents (mol%)	Fluctuation coefficient	Sensiti- zing dye
EM7	Cubic	1.1	99.0	0.1	(ExS-4)
EM8	Cubic	0.8	99.0	0.1	(ExS-4)
EM9	Cubic	0.45	98.5	0.09	(ExS-3,-5)
EM10	Cubic	0.34	98.5	0.09	(ExS-3,-5)
EM11	Cubic	0.45	98.5	0.09	(ExS-8,-12)
EM12	Cubic	0.34	98.4	0.10	(ExS-8,-12)

Then, Samples (D-2)-(D-16) were prepared by exchanging the magenta coupler in the third layer (green-sensitive layer) of Sample (D-1) with another magenta coupler of equal mol and/or by adding a preservability improving compound of the present invention. The details of the Samples (D-2)-(D-16) are shown in Table 5.

Each thus prepared sample was subjected to an exposure through an optical wedge and then to the processing procedure (IV) described below to obtain a color image.

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Processing Procedure(VI)

Step	Temperature (°C)	Time (sec.)	Replenisher Amount (ml/m²)*	Tank Capacity (1)
Color developing	35	45	161	17
Bleach-fixing	30-36	45	215	17
Stabilizing ①	30-37	20	· _	10
Stabilizing ②	30-37	20	-	10
Stabilizing ③	30-37	20	-	10
Stabilizing 4	30-37	30	248	10
Drying	70-85	60		

 $^{\ ^*}$ per m^2 of the photographic material

Stabilizing steps were carried out in a four-tank counter-current mode from tank of stabilizing 1 toward tank of stabilizing 1.

The composition of each processing solution was as follows:

Color Developing Solution

45		Tank Solution	Replenisher
	Water	800 ml	800 ml
	Ethylenediaminetetraacetic acid	2.0 g	2.0 g

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	5,6-dihydroxybenzene-1,2,4-		
5	trisulfonate	0.3 g	0.3 g
	Triethanolamine	8.0 g	8.0 g
10	Sodium chloride	1.4 g	_
10	Potassium carbonate	25 g	25 g
	N-Ethyl-N-(β-methanesulfonamidoe	thyl)-	
15	3-methyl-4-aminoaniline sulfonate	5.0 g	7.0 g
	Diethylhydroxylamine	4.2 g	6.0 g
20	Brightening agent (4,4 -diamino-		
	stilbene series)	2.0 g	2.5 g
	Water to make	1000 m&	1000 mL
25	pH (25°C)	10.05	10.45
30 35	Bleach-fixing Solution (both tank solution and replenisher) Water Ammonium thiosulfate (70% solution) 100 m t Sodium sulfite 17 g Ammonium iron (III) ethylenediaminetetraacetate 55 g Disodium ethylenediaminetetraacetate 5 g Glacial acetic acid 9 g Water to make 1000 m t pH (25°C) 5.40	400 mt	
	Chabittaine Call of the Market Call		

Stabilizing Solution (both tank solution and replenisher) Formalin (37% solution)

Formalin-sulfic acid adduct 0.7 g

5-Chloro-2-methyl-4-isothiazoline-3-one 0.02 g

2-Methyl-4-isothiazoline-3-one 0.01 g

Copper sulfate 0.005 g

Water to make 1000 mt

pH (25 C) 4.0

Then, magenta reflective density (stain) measurements were carried out on the samples as in Example 3 and Example 4, that is, on the processed samples after one hour of the development processing, on the processed samples after being kept for 14 days at 60°C and 70% RH, and on the processed samples after being kept for 100 days at room temperature. The values of increments of magenta stain after one hour were evaluated. The results are shown in Table 5.

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			·																
5 -		Remarks	Comparative	Example	=	=	; ;	Ξ	This	Invention Comparative	Example .	This	Invention Comparative	Example "	This	Invention Comparative	Example"	This Invention	
10	genta Stain	100 days at R.T.	η0.0	0.04	0.02	40.0	0.02	0.01	0.00	0.02	0.02	0.00	0.01	0.01	0.00	0.02	0.02	00.00	
20	Increment of Magenta	14 days at 60°C, 70%RH	0.05	0.04	0.03	0.04	0.02	0.02	0.01	0.02	0.02	0.01	0.02	0.02	0.00	0.02	0.02	0.01	
25 9 1 Q 8 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Amount of Additive	ol% to coupler)					, 10	10	ιυ +	10	10	5 + 5	10	10	r. 10.	10	10,	ν. •	
35	Additive						(1-18)	(61-111)	(61-III) + (81-I)	(1-26)	(111-1)	(I-26) + (III-7)	(11-3)	(h-III)	(II-3) + (III-4)	(I-5)	(111-19)	(I-5) + (III-19)	
45	Magenta	Coupler	ExM-1	ExM-2	ExM-3	ExM-4	ExM-1	=	:	ExM-2	5	=	ExM-3	=	=	ExM-4	=	=	
50	Sample		D - 1	D - 2	D - 3	۵ - د	0 - 5	9 - Q	L - 0	B - Q	6 - Q	D -10	D -11	D -12	ņ -13	D -14	D -15	D -16	

As is apparent from the results in Table 5, as in the above-described Examples, magenta stain increased on each comparative sample, but the increment of stain was not substantially or at all observed on each sample that used the preservability improving compounds (A) and (B) of the present invention in combination, whereas the stain of samples that used compounds (A) or (B) individually were not adequately prevented.

Example 6

Photographic samples for comparison (A-1)-(A-3) prepared in Example 1 were respectively subjected to an exposure through an optical wedge and then to processing according to a comparative procedure (VII) and to processing according to the present invention (VIII) and (IX) to obtain color images.

Processing Procedure(VIII)

Step	Temperature (°C)	Time
Color developing	38	1 min. 40 sec.
Bleach-fixing	30-34	l min.
Rinsing ①	30-34	20 sec.
Rinsing ②	30-34	20 sec.
Rinsing ③	30-34	20 sec.
Drying	70-80	50 sec.

Rinsing steps were carried out in a three-tank countercurrent mode from tank of rinsing ③ toward tank of rinsing ①.

The composition of the processing solutions were as follows:

```
Color Developing SolutionWater
                                   800 m t
Diethylenetriaminepentaacetate
1-Hydroxyethylidene-1,1-diphosphonate (60%)
                                                 2.0 g
Nitolirotriacetic acid
                       2.0 g
Benzyl alcohol
                  16 m t
Diethylene glycol
                    10 mt
Sodium sulfite
                 2.0 g
Potassium bromide
                      0.5 g
Potassium carbonate
                        30 g
N-Ethyl-N-(\beta-methanesulfonamidoethyl)-3-methyl-4-aminoaniline sulfonate
                                                                           5.5 g
Brightening agent (WHITEX4B, made by Sumitomo Chemical)
Water to make
                  1000 mt
     pH (25°C)
                    10.25
```

Bleach-fixing SolutionWater 400 mt
Ammonium thiosulfate (70%) 200 mt
Sodium sulfite 20 g
Ammonium iron (III) ethylenediaminetetraacetate 60 g
Disodium ethylenediaminetetraacetate 10 g
Water to make 1000 mt
pH (25°C) 7.00

Hinsing SolutionBenzotriazole 1.0 g
Ethylenediamine-N,N,N',N'-tetramethylenephosphonate 0.3 g
Water to make 1000 m t
pH (25°C) 7.50

<u>Processing Procedure(VIII)</u> (the present invention) The same as Processing Procedure (VII), except that 10 g of the exemplified compound (III-10) is contained in the rinsing solution.

Processing Procedure(IX) (the present invention)

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The same as processing procedure (VII), except that 10 g of the exemplified compound (III-17) is contained in the rinsing solution.

Then, as in Example 3, magenta reflective density (stain) was measured at a non-image area of each sample at the point of one hour after the development process. The same stain measurements were carried out again on the processed samples after being kept for 14 days at 60°C and 70% RH, and on the processed samples after being kept for 100 days at room temperature. The increments of magenta stain after one hour for the samples were evaluated (Table 6).

									٠,			
		Remerks	Comparative	n Exampte	=	ε	This	"	Comparative	This		
	lagenta Stain	100 days at R.T.	0.15	0.07	0.05	0.05	0.01	0.01	90.0	10.0	0.01	
	Increment of Magenta Stain	14. days at 60°C, 70%RH	0.16	60.0	90.0	0.08	0.01	0.02	60.0	0.01	0.01	
· • • • • • • • • • • • • • • • • • • •	Processing	Procedure	(AII)	(VII)	(VII)	(VIII)	(IIIA),	(VIII)	(XI)	(XI)	(XI)	
Table 6	Amount of	Additive (molf to coupler)	-	20	20 }		20	20	1	20	20	
	Additive	(Exemplified Compound)		(I-4)	(1-27)		(1-1)	(I-27)		(1-1)	(1-27)	
	Magenta	Coupler	м - 30	ŧ	=		=	=	=	=	E	
· ·	Sample		A - 1	A - 2	A - 3	A - 1	A - 2	A - 3	A - 1	A - 2	A - 3	

As is apparent from the results in Table 6, with the inclusion of the preservability improving compounds in the photographic materials after the developing steps, the increments of magenta stain are lowest on the samples in which both of the preservability improving compounds (A) and (B) were applied to.

Example 7

The preparation procedures of the photographic materials in Examples 1-6 were repeated, except that the cyan couplers were changed to (ExC-1)-(ExC-6), respectively. The thus prepared samples were evaluated for magenta stain in the same manner as Examples 1-6. From the results of the evaluation, it is clear that the most preferable prevention of stain was obtained with the combined use of the preservability improving compounds (A) and (B), as in the above-described Examples.

As is evident from these results, the stain increment with a lapse of time is remarkably prevented by the practice of the present invention, and this excellent effect is maintained even if the structure of photographic materials and processing procedures are varied.

Example 8

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Samples (C-17) to (C-24) were prepared by repeating the preparation procedures of Sample (C-1) in Example 3, except the changing of the yellow coupler (equal mole exchanging) and the solvent in the first layer (blue-sensitive emulsion layer), with or without adding the preservability improving compounds of the present invention; changing the magenta coupler (equal mole exchanging) and the solvent in the third layer (green-sensitive emulsion layer), with or without adding the preservability improving compounds of the present invention; and changing the cyan coupler (equal mole exchanging) and the solvent in the fifth layer (red-sensitive emulsion layer), with or without adding the preservability improving compounds of the present invention. The details of the exchanged compositions are shown in Table 7.

Table 7

Sample	Main component	First Layer	Third Layer	Pifth Layer
(0-17)	Coupler Preservability improving compound Solvent	ExY-3. - Solv-8	ExM-4 ° . - Solv-1	ExC-4/ExC-8 = 1/1
(C-18)	Coupler Preservability improving compound Solvent	Same as (C-17) I-32 (5 mol%)* Same as (C-17)	Same as (C-17) I-74 (20 molf)* Same as (C-17)	Same as (C-17) I-82 (5 mol#)* Same as (C-17)
(c-19)	Coupler Preservability improving compound Solvent	Same as (C-18) Same as (C-18) Same as '(C-18)	Same as (C-17) III-30 (20 mol\$)* Same as (C-17)	Same as (C-18) Same as (C-18) Same as (C-18)
(0-50)	Coupler Preservability improving compound Solvent	Same as (C-18) Same as (C-18) Same as (C-18)	Same as (C-17) I-74(10 mol%)*, III-30(10 mol%)* Same as (C-18) Same as (C-17)	Same as (C-18) Same as (C-18) Same as (C-18)
(C-21)	Coupler Preservability improving compound Solvent	ExY-1/ExY-2 = 1/1 - Solv-8	ExM-2 I-70 (20 mol#)* Solv-4/Solv-6 = 1/2	ExC-3 - Solv-8
(C-22)	Coupler Preservability improving compound Solvent	Same as (C-21.) Same as (C-21.) Same as (C-21.)	Same as (C-21) III-40 (20 molf)* Same as (C-21)	Same as (C-21) Same as (C-21) Same as (C-21)
(0-23)	Coupler Preservability improving compound Solvent	Seme as (C-21) Same as (C-21) Same as (C-21)	Same as (C-21) I-70(10 molx)*,III-40(10 molx)* Same as (C-21)	Same as (C-21) Same as (C-21) Same as (C-21)
(C-24)	Coupler Preservability improving compound Solvent	Same as (C-21) I-71 (5 molg)* Same as (C-21)	Same as (C-21) I-57(10 mol%)*, III-30(10 mol%)* Same as (C-21)	Same as (C-21) I-71 (5 mol#)* Same as (C-21)

Note) *: molf based on coupler

Each of the thus prepared samples was subjected to an exposure through an optical wedge and then processed according to the processing procedure (IV) shown in Example 3.

Then, reflective densities of yellow, magenta, and cyan at a non-image area of each processed sample (C-17) to (C-24) were measured at the point of one hour after development processing, and again after being kept for 14 days at 60°C and 70% RH. The results are shown in Table 8 as values of increments of stain after one hour concerning yellow, magenta, and cyan reflective densities.

Table-8

Sample	Increment of being kept 60°C 70% RH	Remarks					
·	Yellow(D _B)	$Magenta(D_G)$	Cyan(D _C)				
(C-17)	0.15	0.11	0.08	Comparative Example			
(C-18)	0.05	0.05	0.03	n Bygmbre			
(C-19)	0.06	0.04	0.05	This Invention			
(C-20)	0.02	0.01	0.02	***			
(C-21)	0.14	0.10	0.15	Comparative			
(C-22)	0.11	0.05	0.11	Example "			
(C-23)	0.07	0.02	0.10	This Invention			
(C-24)	0.02	0.01	0.02	11			

As is apparent from the results of Table 8, yellow stain, magenta stain and cyan stain increase remarkably on the samples (C-17) and (C-21) that did not use the preservability improving compounds in the processing procedure (IV), but the increments of stain are prevented considerably by adding the preservability improving compound (A) into each layer [Sample (C-18)]. Further better effect is obtained by adding the preservability improving compound (B) [Sample (C-19)]. It is noted that the increments of stain in the first and fifth layers are prevented somewhat by using the preservability improving compound (A) in the third layer [Sample (C-22)]. However, the effect of the preservability improving compound is not sufficient with the individual use of compounds (A) and (B). In contrast, a superior effect is obtained by the combined use of these preservability improving compounds (A) and (B) in the same layer [Samples (C-20), (C-23) and (C-24)]. And also, it is noted that the increment of each stain is almost completely prevented by using a small amount of the preservability improving compound (A) in the processing procedure such as in processing procedure (IV) [Samples (C-20) and (C-24)].

50 Example 9

A color photographic material (H-1) was prepared by multi-coatings composed of the first to the fourteenth layer as hereinbelow described on a both-sides polethylene laminated paper base. A white pigment (TiO₂) and a small amount of bluish dye (ultramarine blue) were included in the first layer side of the polyethylene film laminated.

Composition of photosensitive layers

In the following compositions, each ingredient is indicated in g/m² of a coating amount, but the coating amount of the silver halide is shown in g/m² in terms of silver.

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First layer: Antihalation layer Black colloidal silver 0.10

Gelatin 1.30

10

Second Layer: Intermediate layer Gelatin 0.70

Third layer: Red-sensitive emulsion (low sensitivity) layer Silver bromide emulsion spectral-sensitized by red-sensitizing dye (ExS-1, -2 and -3) (average grain size: 0.3 μm, grain size distribution: 8 %, octahedral)

Silver bromide emulsion spectral-sensitized by red-sensitizing dye (ExS-1, -2 and -3)(average grain size: 0.45 µm, grain size distribution: 10%, octahedral) 0.10

20 Gelatin 1.00

Cyan coupler (ExC-1) 0.14

Cyan coupler (ExC-2) 0.07

Discoloration inhibitor (equal amount mixture of Cpd-2, -4, -5 and -9) 0.12

Dispersion medium for coupler (Cpd-5) 0.20

25 Solvent for coupler (equal amount mixture of Solv-1, -2 and -3) 0.06

Fourth layer: Red-sensitive emulsion (highly sensitive) layer Silver bromide emulsion spectral-sensitized by red-sensitizing dye (ExS-1, -2 and -3) (average grain size: 0.75 µm, grain size distribution: 10%, octahedral) 0.15

Octanoular) U.

Gelatin 1.00

Cyan coupler (ExC-1) 0.20

Cyan coupler (ExC-2) 0.10

Discoloration inhibitor (equal amount mixture of Cpd-2, -3, -4 and -9) 0.15

35 Dispersion medium for coupler (Cpd-5) 0.30

Solvent for coupler (equal amount mixture of Solv-1, -2 and -3) 0.10

Fifth layer: Intermediate layer Gelatin 1.00

40 Color mix inhibitor (Cpd-7) 0.08

Solvent for color mix inhibitor (Solv-4 and -5) 0.16

Polymer latex (Cpd-8) 0.10

45 Sixth layer: Green-sensitive emulsion (low sensitivity)layer Silver bromide emulsion spectral-sensitized by green-sensitizing dye (ExS-3 and -4) (average grain size: 0.28 μm, grain size distribution: 8%, octahedral) 0.04

Silver bromide emulsion spectral-sensitized by green-sensitizing dye (ExS-3 and -4) (average grain size : $0.45 \mu m$, grain size distribution : 10 %, octahedral) 0.06

50 Gelatin 0.80

Magenta coupler (ExM-1) 0.10

Discoloration inhibitor (Cpd-9) 0.10

Stain inhibitor (Cpd-10) 0.01

Stain inhibitor (Cpd-11) 0.001

Stain inhibitor (Cpd-12) 0.01

Dispersion medium for coupler (Cpd-5) 0.05

Solvent for coupler (equal amount mixture of Solv-4 and -6) 0.15

```
Seventh layer: Green-sensitive emulsion (highly sensitive) layer Silver bromide emulsion spectral-sensitized
     by green-sensitizing dye (ExS-3)(average grain size : 0.9 µm, grain size distribution : 10 %, octahedral
     0.10
     Gelatin
                 0.80
     Magenta coupler (ExM-1)
                                   0.10
     Discoloration inhibitor (Cpd-9)
                                       0.10
     Stain inhibitor (Cpd-10)
                                 0.10
     Stain inhibitor (Cpd-11)
                                 0.001
     Stain inhibitor (Cpd-12)
                                 0.01
     Dispersion medium for coupler (Cpd-5)
     Solvent for coupler (equal amount mixture of Solv-4 and -6)
                                                                     0.15
     Eighth layer: Intermediate layer Same as the fifth layer.
 15
     Ninth layer: Yellow filter layer Yellow colloidal silver
                                                              0.20
     Gelatin
                1.00
     Color mix inhibitor (Cpd-7)
     Solvent for color mix inhibitor (equal amount mixture of Solv-4 and -5)
                                                                                0.15
     Polymer latex (Cpd-8)
                               0.10
     Tenth layer: Intermediate layer Same as the fifth layer.
25
     Eleventh layer: Blue-sensitive emulsion (low sensitivity) layer Silver bromide emulsion spectral-sensitized by
     blue-sensitizing dye (ExS-5)(average grain size: 0.35 µm, grain size distribution: 8%, tetradecahedral)
    Silver bromide emulsion spectral-sensitized by blue-sensitizing dye (ExS-5)(average grain size : 0.45 µm,
     grain size distribution: 10%, tetradecahedral)
                                                       0.10
     Gelatin
                0.50
     Yellow coupler (ExY-1)
                                0.20
    Stain inhibitor (Cpd-11)
                                0.001
    Discoloration inhibitor (Cpd-6)
    Dispersion medium for coupler (Cpd-5)
                                                0.05
    Solvent for coupler (Solv-2)
                                    0.05
    Twelfth layer: Blue-sensitive emulsion (highly sensitive) layer Silver bromide emulsion spectral-sensitized
    by blue-sensitizing dye (ExS-5 and -6) (average grain size : 1.2 µm, grain size distribution : 10 %,
    tetradecahedral)
                         0.25
    Gelatin
                1.00
    Yellow coupler (ExY-1)
                               0.40
    Stain inhibitor (Cpd-11)
                                0.002
    Discoloration inhibitor (Cpd-6)
    Dispersion medium for coupler (Cpd-5)
                                                0.05
    Solvent for coupler (Solv-2)
50
    Thirteenth layer: UV absorbing layer Gelatin
    UV absorbent (equal amounts mixture of Cpd-1, -3 and -13)
                                                                    1.00
    Color mix inhibitor (equal amount mixture of Cpd-6 and -14)
                                                                    0.06
    Dispersion medium (Cpd-5)
                                    0.20
    Solvent for UV absorbent (equal amount mixture of Solv-1 and -2)
                                                                          0.15
```

0.02

0.02

Irradiation inhibitor dye (equal amount mixture of Cpd-15 and -16)

Irradiation inhibitor dye (equal amount mixture of Cpd-17 and -18)

0 277 589

Fourteenth layer: Protective layer Fine grain size silver chlorobromide emulsion (silver chloride: 97 mol%,

average grain size : 0.2 µm) 0.15 Modified polyvinyl alcohol 0.02

Gelatin 1.50

5 Gelatin hardner (H-1) 0.17

Next, the preparation procedure of the emulsion for the respective layers, except the fourteenth layer, is exemplified as follows:

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Preparation of emulsion

An aqueous solution containing potassium bromide and silver nitrate was added to an aqueous solution of gelatin containing 0.3 g/moleAg of 3,4-dimethyl-1,3-thiazoline-2-thione with vigorous agitation at 75°C over about 20 min, to obtain a mondisperse silver bromide emulsion of octahedral cystalline particles having an average grain size of 0.40 μ m. A chemical sensitizing treatment of the thus obtained emulsion was carried out by adding 6 mg/moleAg of sodium thiosulfate and 7 mg/moleAg of chloroauric acid (tetrahydrate) and heating it at 75°C for 80 min. Thus obtained silver bromide grains were bought up as a core in the same precipitating conditions as the first precipitating process to obtain finally a monodisperse core-shell silver bromide emulsion of octahedral shaped grains having an average grain size of 0.7 μ m. The fluctuation coefficient of the grain size distribution of this emulsion was about 10%.

A further chemical sensitization of this emulsion was carried out by adding 1.5 mg/mol•Ag of sodium thiosulfate and 1.5 mg/mol•Ag of chloroauric acid (tetrahydrate) and heating it at 60°C for 60 min, to obtain an internal latent-image type silver halide emulsion.

Further, 10 ³ weight % of the compound (N-1) to the coating amount of silver halide and 10 weight % of the compound (ExZS-1) were included in each layer as a nucleating agent and nucleation accelerator, respectively.

In addition, the same auxiliary agents for emulsification and dispersion and coating aids as in Example 2 were used. As the stabilizer in the layers containing silver halide or colloidal silver, compound (Cpd-19), (Cpd-20), and (Cpd-21) were used.

The compounds used in the Examples were as follows:

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ExS-1

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$$\begin{array}{c|c}
S & C_2H_5 \\
+ & CH = C - CH = \\
N & CH_2)_3SO_3^{-1} & CH_2)_3SO_3H
\end{array}$$

ExS-2

20
$$C\ell \xrightarrow{S} C_{2}H_{5}$$

$$CH = C - CH = C$$

$$CH_{2})_{3}SO_{3}^{-} (CH_{2})_{3}SO_{3}H$$

ExS-3

ExS-4

$$\begin{array}{c|c}
C_2H_5 \\
C_2H_4SO_3
\end{array}$$

$$\begin{array}{c|c}
C_2H_5 \\
C_2H_4SO_3
\end{array}$$

$$\begin{array}{c|c}
C_2H_4SO_3H
\end{array}$$

ExS-5

Cpd-1

$$\begin{array}{c|c}
 & \text{HO} & \text{C}_4\text{H}_9 \text{ (sec)} \\
 & & \\
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Cpd-2

Cpd-3

Cpd-4

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HO
$$C_4H_9(t)$$
 $C_4H_9(t)$ $C_4H_9(t)$

Cpd-5

25 Cpd-6

$$\begin{bmatrix} C_4H_9(t) \\ HO - CH_2 \\ C_4H_9(t) \end{bmatrix}_2 - C - \begin{bmatrix} CH_3 \\ CH_3 \\ CH_3 \end{bmatrix}_2$$

$$CH_3 CH_3 \\ CH_3 CH_3 \\ CH_3 CH_3 CH_3 CH_3$$

35 Cpd-7

Cpd-8

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Polyethyl acrylate

Cpd-9

Cpd-10

 $C_5H_{11}(t)$

Cpd-11

$$C_{pd}-12 \qquad C_{5}H_{11}(t)$$

$$CONHC_{3}H_{6}O - C_{5}H_{11}(t)$$

$$CONHC_{3}H_{6}O - C_{5}H_{11}(t)$$

Cpd-13

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 $\begin{array}{c|c} C\ell & HO & C_4H_7(t) \\ \hline & N & O \\ \hline & CH_2CH_2COC_8H_{17} \end{array}$

Cpd-14

OH C₈H₁₇ (sec) OH

Cpd-15

C₂H₅OCO CH-CH=CH CO₂C₂H₅
N
O
HO
N
SO₃K
SO₃K

Cpd-16

 C_2H_5OCO CH-CH=CH

N

O

HO

N

CO₂C₂H₅

CO₂C₂H₅

CH₂)₃SO₃K

(CH₂)₃SO₃K

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Cpd-17

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C₂H₅OCO CH-CH=CH-CH=CH
$$\frac{1}{N}$$
 CO₂C₂H₁
NO
HO
N
SO₃K
SO₃K

Cpd-18

Cpd-19

Cpd-20

5

70

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25

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Cpd-21

20 EXC-1

$$C_{2}H_{5} \xrightarrow{OH} C_{4}H_{9}$$

$$C_{2}H_{5} \xrightarrow{I} C_{5}H_{11}(t)$$

$$C_{5}H_{11}(t)$$

EXC-2

(t)
$$C_5H_{11}$$
 OCHCONH CL

ExM-I

ExY-1

5 $(CH_3)_3CCOCHCONH$ O = N O = N O = N O = N + COCHO $(t) C_5 H_{11}$ $C_2 H_5$

Solv-1 Di(2-ethylhexyl)phthalate

Solv-2 Trinonylphosphate

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Solv-3 Di(3-methylhexyl)phthalate

Solv-4 Tricrezylphosphate

Solv-5 Dibutylphthalate

Solv-6 Trioctylphophate

Solv-7 Dioctylsebacate

H-1 1,2-bis(vinylsulfonylacetoamido)ethane

$$(N-I-9)$$

SH

NNNN

CONH

CH2CECH-CLO4

(ExZS-1)

HS S S-(CH₂)₆-N
$$CH_3$$
-HCL

Then, as shown in Table 9, Samples (H-2) to (H-14) were prepared by repeating the preparation procedures of Sample (H-1) except the changing of the magenta couplers and the preservability improving compounds (Cpd-10) and (Cpd-12) in the sixth layer and the seventh layer respectively.

Each of the thus prepared samples was subjected to an exposure through an optical wedge and then to a color development process according to the following processing procedure (X).

0 277 589

Processing Procedure(X)

Step	Time (sec.)	Temperature (°C)
Color develop	ing 90	. 38
Bleach-fixing	45	38
Water washing	① 45	38
Water washing	2 45	38

The water washing steps were carried out by a so-called countercurrent replenishing mode, in which the replenisher is fed to bath of water washing ②, and the overflow water from bath of water washing ③ is fed to bath of water washing ①.

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Color Developing Solution

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10		Mother solution
	Diethylenetriaminepentaacetic acid	0.5 g
15	1-Hydroxyethylidene-1,1-diphosphonate	0.5 g
	Diethylene glycohol	8.0 g
20	Benzyl alcohol	12.0 g
	Sodium bromide	0.7 g
	Sodium sulfite	2.0 g
25	N,N-diethylhydroxylamine	3.5 g
	Triethylenediamine (1,4-diazabicyclo [2	2,2,2]
30	octane	3.5 g
	3-Methyl-4-amino-N-ethyl-(β-methanesulf	Conamido-
	ethyl)-aniline	6.0 g
35	Potassium carbonate	30.0 g
	Brightening agent (Stilbene series)	1.0 g
40	Water to make	1000 m &
	pH (adjusted with potassium hydroxide o	r hydrochloric
	acid)	10.50
45	Bleach-fixing Solution	× ·
		Mother solution
50	Ammonium thiosulfate	110 g
	Sodium hydrosulfite	14.0 g

Ammonium iron(III) ethylenediaminetetra
acetate dihydrate - 40.0 g

Disodium ethylenediaminetetraacetate

dihydrate 4.0 g

Water to make 1000 ml

pH (adjusted with aqueous ammonia or hydrochloric

acid) 10.50

Washing Water

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Purified water (de-ionized tap water by ion-exchange treatment, containing under 1 ppm of all cations except the hydrogren ion and all anions except the hydroxide ion)

Then, magenta reflective density (stain) was measured at a non-image area of each sample at the point of one hour after the development processing. The same stain measurements were carried out again on the processed samples after being kept for 6 days at 80 C and 70% RH, and on the processed samples after being kept for 100 days at room temperature. The increments of magenta stain to that of one hour after processing for each sample are shown in Table 9.

				Table 9		5 0 0	Motes
Башр 1 е	Magenta Coupler		Freservability improving Compound	Addition (molf)*	at 80°C, 70%RH at R.T.	IOO days	P 200
(H-1)	ExM-1 (M-26)	(A)	: Cpd-10 (I-32) : Cpd-12 (III-30)	10	0.01	0.00	This Invention
(H-2)	=	(A)	1 1	1 1	0.15	0.03	Comparative Example
(н-3)	E	(B)	: Cpd-10 (I-32)	, 20	0.05	0.03	1
(H-4)	· =	(A)	; Cpd-12 (III-30)	20.	60.0	0.03	=
(H-5)	F	(A)	(1- <u>7</u> 0)	20	90.0	0.03	=
(9-н)	Ε	(A) (B)	: (IIÎ-40)	20	0.08	0.02	1
(H-7)	E	(A)	(I-70) (III-40)	10	. 0.01	0.01	This Invention
(н-8)	(M-30)	(A) (B)	1 1	, ,	0.16	0.13	Comparative Example
(н-9)	t	(A) (B)	(1-57)	0.2	90°0 .	0.03	a
(H-10)	±	(A)	(I-57) (III-30)	22	0.01	00.0	This Invention
(H-11)	(M-37)	: (¥)	1 1		0.16	0.10	Comparative Example
(H-12)	11	(A):	(I-74) (III-32)	10	0.01	0.01	This Invention
(н-13)	(M-43)	(A): (B):			0.05	0.04	Comparative Example
(H-14)	=	(A)	(I-58) (TII-36)	10	0.01	0.00	This Invention

Note) *: mol% on coupler

As is apparent from the results of Table 9, the stain increments over a lapse of time on the processed photographic material were prevented remarkably by using in combination the preservability improving compounds (A) and (B) of the present invention.

Further, even when the ratio of the silver bromide emulsion to the silver chlorobromide emulsion is varied (in the range that silver chloride is 0.5-99.5 mol%), nearly the same effects as in Table 9 were attained.

Having described our invention as related to the embodiment, it is our intention that the invention be not limited by any of the details of the description, unless otherwise specified, but rather be construed broadly within its spirit and scope as set out in the accompanying claims.

Claims

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1. A silver halide color photographic material containing both a compound (A), that combines chemically with the aromatic amine developing agent remaining after a color development processing to produce a chemically inactive and substantially colorless compound, and a compound (B), that combines chemically with the oxidized product of the aromatic amine developing agent remaining after the color development processing to produce a chemically inactive and substantially colorless compound.

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- 2. The color photographic material as claimed in claim 1, wherein compound (A) is selected from compounds that can react with a rate constant k_2 (at 80°C) of the secondary reaction with p-anisidine within the range of 1.0 t/mol.sec to 1 x 10 5 1 /mol.sec.
- 3. The silver halide color photographic material as claimed in claim 1, wherein compound (B) is selected from compounds having a nucleophilic group derived from a nucleophilic functional group that has a Pearson's nucleophilic "CH₃I value of 5 or greater.
- 4. The silver halide color photographic material as claimed in claim 1, wherein the amount of compound (A) or (B) is 1 x 10 2 to 10 mol per mol of a coupler employed.
- 5. The silver halide color photographic material as claimed in claim 1, wherein the amount of compound (B) contained is in the range of 2×10^{-2} to 2×10^{-1} per mol of a coupler employed.
- 6. The silver halide color photographic material as claimed in claim 1, wherein compound (A) or (B) is co-emulsified with a coupler, the oil/coupler weight ratio being from 0.01 to 2.0.
- 7. The silver halide color photographic material as claimed in claim 1, wherein at least one of the couplers represented by the following formulae is employed:

General formula (Y-I)

wherein R₁₁ represents a substituted or unsubstituted N-phenylcarbamoyl group, and Z₁₁ represents a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent;

General formula (Y-II)

$$(R_{12})_s$$
 $COCH-R_{11}$

wherein R₁₁ represents a substituted or unsubstituted N-phenylcarbamoyl group, Z₁₁ represents a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent, R₁₂ represents a hydrogen atom or a substituent group, and s is an integer of 1 to 5;

General formula (M-I)

wherein R₂₁ represents an alkyl group, an aryl group, an acyl group, or a carbamoyl group; Ar represents a phenyl group or a phenyl group substituted by one or more halogen atoms, alkyl groups, cyano groups, alkoxy groups, alkoxycarbonyl groups, or acylamino groups; and Z₂₁ represents a hydrogen atom or a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent;

General formula (M-II)

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$$R_{22}$$
 Z_{21}
 N
 Z_{24}
 Z_{22}
 Z_{23}

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wherein R_{22} represents a hydrogen atom or a substituent group; Z_{21} represents a hydrogen atom or a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent; Z_{22} ,

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 Z_{23} and R_{24} represent - C = 1, -N = 1 or -NH = 1; between the bonds of $Z_{24} = Z_{23}$ and $Z_{25} = Z_{25}$, one is a double bond and the other is a single bond; and when the $Z_{25} = Z_{25}$ is a carbon-carbon double bond, the double bond may be part of an aromatic ring;

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General formula (C-I)

OH
$$R_{33} \longrightarrow NHCOR_{3}$$

$$R_{32} \longrightarrow Z_{31}$$

wherein R₃₁ represents an alkyl group, a cycloalkyl group, an aryl group, an amino group, or a heterocyclic group; R₃₂ represents an acylamino group or an alkyl group; R₃₃ represents a hydrogen atom, an alkyl group or an alkoxy group; R₃₃ and R₃₂ may bond together to form a ring; and Z₃₁ represents a hydrogen atom, a halogen atom, or a group that can split off in the reaction with the oxidized product of the aromatic primary amine color developing agent;

so and

the above couplers may form a dimer or even higher polymer.

- 8. The silver halide color photographic material as claimed in claim 1, wherein the silver halide comprises silver chloride, silver bromide, or a mixed silver halide.
- 9. The silver halide color photographic material as claimed in claim 1, wherein compound (A) is represented by the following general formula (I) or (II):

General formula (i) R₁ + A + X

General formula (II) R₂-C = Y

wherein R_1 and R_2 each represent an aliphatic group, an aromatic group, or a heterocyclic group; X represents a group that can react with the aromatic amine developing agent to cause splitting-off; A represents a group that can react with the aromatic amine developing agent to form a chemical bond; n is 1 or 0; B represents a hydrogen atom, an aliphatic group, an aromatic group, a heterocyclic group, an acyl group, or a sulfonyl group; Y represents a group that can facilitate the addition of the aromatic amine developing agent to a compound having general formula (II), and R_1 and X together or Y and R_2 or B together may combine to form a ring structure.

10. The silver halide color photographic material as claimed in claim 9, wherein the aliphatic group of R₁, R₂ and B represents a straight chain, branched chain or cyclic alkyl groups, alkenyl group or alkynyl group; the aromatic group of R₁, R₂ and B represents a carbocyclic aromatic and the heterocyclic aromatic group; and the heterocyclic group R₁, R₂ and B represents a 3 to 10-membered heterocyclic group comprising carbon atoms, oxygen atoms, nitrogen atoms, sulfur atoms, or hydrogen atoms;

X represents a group that attaches to A via an oxygen atom, a sulfur atom, a nitrogen atom, or a halogen atom, wherein when X is a halogen atom, n is 0;

A represents a group containing a low electron density atom; and

Y is an oxygen atom, a sulfur atom, = N-R4 or



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wherein R4, R5 and R6 each represent a hydrogen atom, an aliphatic group, an aromatic group, a heterocyclic group, an acyl group, or a sulfonyl group, and R6 and R6 may bond together to form a ring structure.

11. The silver halide color photographic material as claimed in claim 1, wherein compound (B) is represented by the following general formula (III):

General formula (III) R7-Z•M

wherein R₇ represents an aliphatic group, an aromatic group, or a heterocyclic group, Z represents a nucleophilic group, and M represents a hydrogen atom, a metal cation, an ammonium cation, or a protective group.

12. The silver halide color photographic material as claimed in claim 11, wherein the aliphatic group represented by R₇ is a straight chain, branched chain, or cyclic alkyl, alkenyl or alkynyl group; the aromatic group represented by R₇ may be any of a carbocyclic aromatic group and a heterocyclic aromatic group; the heterocyclic group represented by R₇ has a a3 to 10-membered ring structure comprising carbon atoms, oxygen atom, nitrogen atoms, sulfur atoms, or hydrogen atoms;

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Z represents a nucleophilic group having an oxygen atom, a sulfur atom, or a nitrogen atom to chemically combine with the oxidized product of the aromatic amine developing agent; and

M represents a hydrogen atom, a metal cation, an ammonium cation, or a protective group.

13. The silver halide color photographic material as claimed in claim 11, wherein compound (B) is repesented by the following general formula (IV):

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General formula (IV)

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wherein M' represents an atom or an atomic group forming an inorganic or organic salt,

in which $R_{\rm E}$ and $R_{\rm 16}$, which may be the same or different, each represent a hydrogen atom, an aliphatic group, an aromatic group, or a heterocyclic group, or $R_{\rm 15}$ and $R_{\rm 16}$ may bond together to form a 5 to 7-membered ring; $R_{\rm 17}$, $R_{\rm 18}$, $R_{\rm 20}$ and $R_{\rm 21}$, which may be the same or different, each represent a hydrogen atom, an aliphatic group, an aromatic group, a heterocyclic group, an acyl group, an alkoxycarbonyl group, a sulfonyl group, a ureido group, or a urethane group, provided that at least one of $R_{\rm 17}$ and $R_{\rm 18}$ and at least one of $R_{\rm 20}$ and $R_{\rm 21}$ are hydrogen atoms; $R_{\rm 19}$ and $R_{\rm 22}$ represent a hydrogen atom, an aliphatic group, an aromatic group, or a heterocyclic group; $R_{\rm 22}$ further represents an alkylamino group, an arylamino group, an alkoxy group, an aryloxy group, an acyl group, and alkoxycarbonyl group, or an aryloxycarbonyl group; at least two of $R_{\rm 17}$, $R_{\rm 18}$ and $R_{\rm 27}$ may bond together to form a 5 to 7-membered ring;

R₁₀, R₁₁, R₁₂, R₁₃ and R₁₄, which may be the same or different, each represent a hydrogen atom, an aliphatic group, an aromatic group, a heterocyclic group, a halogen atom, -SR₈, -OR₈, - N - R₈, an acyl group, an alkoxycarbonyl

group, an aryloxycarbonyl group, a sulfonyl group, a sulfonamido group, a sulfamoyl group, a ureido group, a urethane group, a carbamoyl group, a sulfo group, a carboxyl group, a nitro group, a cyano group, an alkoxalyl, an aryloxalyl group, a sulfonyloxy group,

- -P(R₆)₂, -P (R₆)₂, -P(OR₆)₃ or a formyl group, wherein R₆ and R₉ each represent a hydrogen atom, an aliphatic group, an alkoxy group, or an aromatic group.
- 14. The silver halide color photographic material as claimed in claim 13, wherein the total of the Hammett's sigma values of R₁₀, R₁₁, R₁₂ and R₁₄ with respect to -SO₂M' is 0.5 or greater.
- 15. The silver halide color photographic material as claimed in claim 1, wherein compound (A) and compound (B) are contained in a layer of the hydrophilic colloid layers on the base.
- 16. A process for preparing a color photograph which comprises processing a silver halide color photographic material in the presence of a compound (A), that combines chemically with the aromatic amine developing agent remaining after a color development processing to produce a chemically inactive and substantially colorless compound, and a compound (B), that combines chemically with the oxidized product of the aromatic amine developing agent remaining after the color development processing to produce a chemically inactive and substantially colorless compound.

- 17. The process for preparing a color photograph as claimed in claim 16, wherein compound (A) and/or compound (B) are contained in one or more layers of the hydrophilic colloid layers on the base of the silver halide color photographic material.
- 18. The process for preparing a color photograph as claimed in claim 16, wherein before, during, or after the color development processing the photographic material is processed with a processing solution to which compound (A) and/or compound (B) has been added, thus allowing them to be contained in the color photograph.
- 19. The process for preparing a color photograph as claimed in claim 16, wherein compound (A) is selected from compounds that can react with a rate constant k_2 (at 80°C) of the secondary reaction with p-anisidine within the range of 1.0 t/mol.sec to 1 x 10 t/mol.sec.
- 20. The process for preparing a color photograph as claimed in claim 16, wherein compound (B) is selected from compounds having a nucleophilic group derived from a nucleophilic functional group that have a Pearson's nucleophilic "CH₃I value of 5 or greater.
- 21. The process for preparing a color photograph as claimed in claim 16, wherein the amount of compound (A) or (B) is 1 x 10 2 to 10 mol per mol of a coupler employed.
- 22. The process for preparing a color photograph as claimed in claim 16, wherein the aromatic amine developing agent is selected from a group consisting of aromatic primary, secondary, and tertiary amine compounds.
- 23. The process for preparing a color photograph as claimed in claim 16, wherein compound (A) is selected from a group consisting of compounds represented by the following general formulae (I) and (II):

General formula (I)
$$R_1 + A + A + A \times X$$

General formula (II) Rz- C = Y

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wherein R₁ and R₂ each represent an aliphatic group, an aromatic group, or a heterocyclic group; X represents a group that can react with the aromatic amine developing agent to cause splitting-off; A represents a group that can react with the aromatic amine developing agent to form a chemical bond; n is 1 or 0; B represents a hydrogen atom, an aliphatic group, an aromatic group, a heterocyclic group, an acyl group, or a sulfonyl group; Y represents a group that can facilitate the addition of the aromatic amine developing agent to a compound having general formula (II); and R₁ and X together or Y and R₂ or B together may combine to form a ring structure.

- 24. The process for preparing a color photograph as claimed in claim 16, wherein compound (B) is represented by the following general formula (III):
- wherein R₇ represents an aliphatic group, an aromatic group, or a heterocyclic group; Z represents a nucleophilic group; and M represents a hydrogen atom, a metal cation, an ammonium cation, or a protective group.
- 25. The process for preparing a color photograph as claimed in claim 16, wherein compound (A) or (B) is added into a color developing solution, a bleaching solution, a fixing solution, a washing solution, or a rinsing solution, the concentration of compound (A) or (B) in the processing solution being 10 ⁵ mol/1 to 10 ¹ mol/1.
 - 26. The process for preparing a color photograph as claimed in claim 12, wherein the color developing solution of the color development processing is substantially free from benzyl alcohol.
- 27. A color photograph obtained by the processing of a silver halide color photographic material and improved in preservability which comprises both a compound (A), that combines chemically with an aromatic amine developing agent remaining after a color development processing to produce a chemically inactive and substantially colorless compound, and a compound (B), that combines chemically with the oxidized product of the aromatic amine developing agent remaining after the color development processing to produce a chemically inactive and substantially colorless compound.
- 28. The color photograph as claimed in claim 27, wherein compound (A) is selected from compounds that can react with a rate constant k_2 (at 80°C) of the secondary reaction with p-anisidine within the range of 1.0 t/mol.sec to 1 x 10 5t /mol.sec.

- 29. The color photograph as claimed in claim 27, wherein compound (B) is selected from compounds having a nucleophilic group derived from a nucleophilic functional group that have a Pearson's nucleophilic "CH₃ I value of s5 or greater.
- 30. The color photograph as claimed in claim 27, wherein the amount of compound (A) or (B) is 1 \times 10 2 to 10 mol per mol of a coupler employed.
- 31. The color photograph as claimed in claim 27, wherein compound (A) is selected from a group consisting of compounds represented by the following general formulae (I) and (II):

General formula (II) $R_z - C = Y$

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wherein R_1 and R_2 each represent an aliphatic group, an aromatic group, or a heterocyclic group; X represents a group that can react with the aromatic amine developing agent to cause splitting-off; A represents a group that can react with the aromatic amine developing agent to form a chemical bond; n is 1 or 0; B represents a hydrogen atom, an aliphatic group, an aromatic group, a heterocyclic group, an acyl group or a sulfonyl group; Y represents a group that can facilitate the addition of the aromatic amine developing agent to a compound having general formula (II); and R_1 and X together or Y and R_2 or B together may combine to form a ring structure.

32. The color photograph as claimed in claim 27, wherein compound (B) is represented by the following general formula (III):

General formula (III) R₇-Z•M

wherein R₇ represents an aliphatic group, an aromatic group, or a heterocyclic group; Z represents a nucleophilic group; and M represents a hydrogen atom, a metal cation, an ammonium cation, or a protective group.

33. The silver halide color photographic material as claimed is claim 1, wherein compound (A) is represented by general formula (I-a), (I-b), (I-c) or (I-d) that can react with the rate constant k_2 (at 80 °C) of the secondary reaction with p-anisidine within the range of 1 x 10 $^{\circ}$ to 1 x 10 $^{\circ}$ t /mol.sec

(I-a)

$$R_{I}$$
-Link-C-O-Ar

(I-b)

$$R_{I} - Link - C - O - C = C$$

$$R_{S} = R_{S}$$

$$R_{S} = R_{S}$$

20 (I-c)

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$$R_1 - Link - C - O - C$$

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(I-d)

$$R_1-Lin_k - C-O-N$$

wherein R₁ represents an aliphatic group, an aromatic group, or a heterocyclic group; Link represents a single bond or -O-; Ar represents an aromatic group, provided that the group released as a result of reaction with an aromatic amine developing agent is not a group useful as a photographic reducing agent; Ra, Rb and Rc, which may be the same or different, each represent a hydrogen atom, an aliphatic, aromatic or heterocyclic group alkoxy group, aryloxy group, heterocycloxy group, alkylthio group, arylthio group, heterocyclothio group, amino group, alkylamino group, acyl group, amido group, sulfonamide group, sulfonyl group, alkoxycarbonyl group, sulfo group, carboxyl group, hydroxyl group, acyloxy group, ureido group, urthane group, carbamoyl group or sulfamoyl group, may combine toghther to form a 5 to 7-membered heterocyclic ring which may be further substituted by a substituent, may form, a spirocyclic ring or bicyclo ring, or may be condensed by an aromatic ring; Z₁ and Z₂ each represent a non-metal atom group necessary to form a 5 to 7-membered heterocyclic ring which may be further substituted by a substituent, may form a spirocyclic ring or bicyclo ring, or may be condensed by an aromatic ring, provided that the compound released as a result of the reaction of Z₁ with an aromatic amine developing agent is not a coupler or 1-phenyl-3-pyrazolidones.

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